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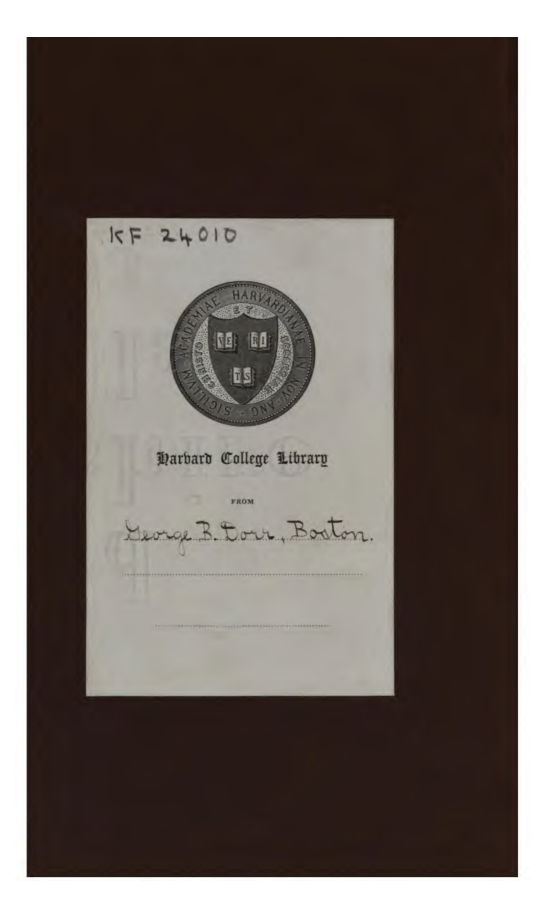
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MANUAL

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OF

DETERMINATIVE MINERALOGY

WITH AN INTRODUCTION

on

BLOW-PIPE ANALYSIS.

,BY

GEORGE J. BRUSH,

PROFESSOR OF MINEBALOGY IN THE SHEFFIELD SCIENTIFIC SCHOOL OF YALE COLLEGE,

> ELEVENTH EDITION. REVISED AND CORRECTED. WITH NEW NOTATION.

NEW YORK: JOHN WILEY & SONS, 15 Aston Place. 1888.

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PREFACE.

The material in this compilation was, for the greater part, prepared almost twenty years since, by Prof. S. W. Johnson and myself as a textbook for the students in our laboratory. Circumstances prevented its publication at that time, but it has served as the basis of a course of lectures and practical exercises annually given in the Sheffield Laboratory.

The plan of instruction has been to have the student work through a course of Qualitative Blowpipe Analysis as introductory to Determinative Mineralogy. For the latter subject we have employed von KOBELL'S *Tafeln Zur Bestimmung der Mineralien*, many of the students taking the work in the original, while others made use of either Erni's or Elderhorst's translations. These "Tables" were translated by Prof. Johnson and myself while we were students of Prof. von Kobell in 1853-4, at Munich, and it was after our suggestion, in 1860, to Prof. Elderhorst, that he introduced von Kobell's "Tables" into the second edition of his "Manual," although he did not avail himself of our translation, which was then offered to him for that purpose.

The "Tables" as now presented are based on the tenth German edition of von Kobell's book. Additions of new species have been made, and in many cases fuller details are given in regard to old species, and the whole material has been thrown into an entirely new shape, which it is believed will greatly facilitate the work of the student. The preparation of the Tables in this form, the idea of which was suggested to me by Prof. W. T. ROEPPER, has been performed, under my supervision, by my assistant Mr. GEORGE W. HAWES, who has also aided me greatly in revising the rest of the work, and in the reading of the proof-sheets.

The main authorities used in the original preparation and later revision of the chapters on Blowpipe Analysis were the works of BERZELIUS and PLATTNER. The third and fourth editions of Plattner, the latter edited by Prof. RICHTER, have been chiefly consulted. The complete work of Plattner, with still later additions by Prof. Richter, has been made accessible to English reading students through an excellent translation by Prof. H. B. CORNWALL, and this cannot be too highly commended to those who desire to become fully acquainted with this important subject.

PREFACE.

In Determinative Mineralogy, besides the works of von Kobell, free use has been made of the treatises of NAUMANN and DANA, especially of the pyrognostic characters contributed by myself to the latter work. This constitutes, in accordance with the original plan of Professor Dana and myself, the Determinative Part of his System of Mineralogy. It is proposed at some future time to add to the volume methods for the determination of minerals by their physical characters.

In conclusion, I take great pleasure in acknowledging my indebtedness to my colleague, Prof. S. W. JOHNSON, who has not only gener usly given me his share in the original work, but has constantly aided me by his advice in the revision here presented.

SHEFFIELD LABORATORY OF YALE COLLEGE, NEW HAVEN, December 15, 1874.

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BLOWPIPE ANALYSIS.

Chapter 1.

APPARATUS AND REAGENTS.

THE MOUTH BLOWPIPE.

1. This little instrument, for centuries employed only by artisans in soldering, and other operations requiring an intense heat, has more recently become an invaluable means of scientific research.*

It is now of the greatest service to the chemist and mineralogist, not only for the recognition of minerals, and the detection of their ingredients, but even for the quantitative separation of several metals from their ores.

The blowpipe serves chiefly for ascertaining the general nature of a body, by revealing some one or more of its ingredients; more rarely it helps to detect all the constituents of a very complex compound, although in but few cases is it possible by its use alone to decide that besides the substances found in a body, no others are present.

The blowpipe enables us in a moment, with no other fuel than that furnished by a common lamp or candle, to produce a most intense heat. In the blowpipe flame not only are most refractory bodies (platinum) melted or volatilized, but the most opposite chemical effects (oxidation and reduction) may be produced. Almost all mineral substances may be made to manifest some characteristic phenomena under its influence, either alone or in presence of certain other substances (reagents), and their nature may be thus surely and easily detected.

2. The Common Blowpipe (Fig. 3) is a conical curved tube of brass, terminating in an orifice as large as a small needle. This simple instrument, when well constructed, answers most ordinary purposes. If used a long time without interruption, the moisture of the breath gathers in drops in the narrow part of the tube, and is finally projected into the flame.

3. In the Chemical Blowpipe a chamber is fixed near the extremity of the instrument which collects the condensed moisture. The most usual form of this

• For a brief history of the use of the blowpipe, see Berzelius' work, translated by J. D. Whitney, Boston, 1845. A more complete history is found in Kopp's Geschichts der Ühemie, II. 44. Braunschweig, 1844. † For Plattner's methods of assaying gold, silver, copper, lead, bismuth, tin, cobalt, nickel

and iron ores, with the help of the blowpipe, see his work cited in the preface.

instrument is shown in Fig. 1 (half size), in which A represents the condenser. Tc admit of emptying this reservoir, it is connected with the tribes by the ground joints b and c. The instrument is also furnished with

a movable jet, a section of which, in correct dimensions, is shown at D This admits of ready cleaning without injury in case of stoppage. Berzelius recommends it to be made from solid platinum, as it then may be easily freed from the soot which is apt to collect upon it, by igniting it in the flame of a spirit lamp, whereby the impurities are burned away. Platinum jets made of foil are too thin at the point, and are thus liable to be easily damaged.

Brass jets are very durable and inexpensive, and may be cleaned, not indeed by heating, but by means of a sharpened splinter of soft wood, which should be introduced for that purpose at the larger end of the jet.

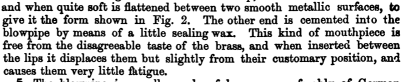
The internal form of the jet is not unimportant. The best shape is that of the section seen in the figure; it is such that the flame produced in using it is always well defined and conical, even when the blast is strong-The jets of the blow-pipe found at the instrumentest. makers' usually need enlarging at the orifice. This is conveniently done with the help of a slender threeedged drill, which may be readily made by grinding down the sides of a large needle.

4. The instrument as shown in Fig. 1, without the trumpet mouthpiece, is of the original form proposed by Gahn, and employed by Berzelius. The beginner is liable to be fatigued in using it, as it requires considerable effort to keep the lips closed about the cylin-drical tube for a long time. Plattner recommends the mouthpiece shown in the figure. It is made of horn or ivory, thirty-five millimetres in its outer diameter, and particular care must be taken that it has the proper curvature, so that in placing it against the lips it may not give an unnecessary or unequal pressure.

A very good mouthpiece may be made from a piece of glass tube, two inches long, and of just such diameter as fits the blowpipe tube. It is strongly and uniformly heated for half its length in the flame of a lamp,



Fig. 1.



5. The blowpipe is usually made of brass, or preferably of German silver. The length of the instrument should be measured by the visual distance of the operator; from seven to nine inches is the ordinary length.

6. In Figs. 3 and 4 is shown how a common blowpipe may be materially improved with but little trouble. A blowpipe being selected that Fig. 3.

gives a good flame, it is cut in two so that the wider part of the tube has a length equal to the visual distance of the operator. The narrow tube is then



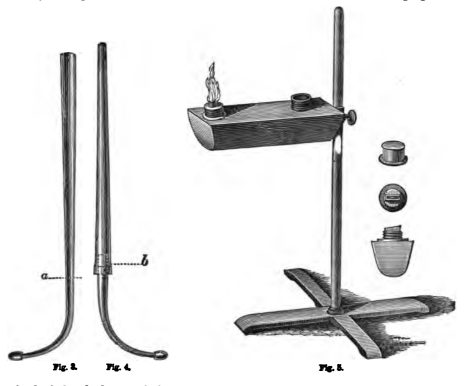
BLOWING.

reversed, and tightly fitted into the wider end of the long tube by means of a perforated cork, thus forming a reservoir for moisture, as seen at a in the figure.

7. Bunsen's gas blowpipe, in which illuminating gas issues from a tubular burner which surrounds the jet of the blowpipe, is sometimes convenient for laboratory use.

BLOWING.

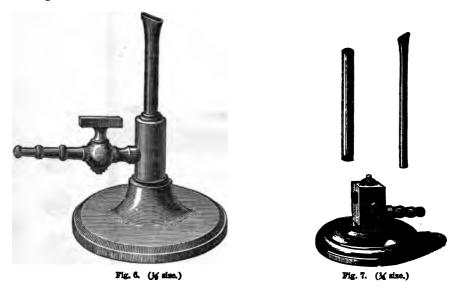
8. In blowpipe operations it is often necessary to maintain an uninterrupted stream of air for several successive minutes. To be able to do this easily, requires some practice. It is best learned by fully distending the cheeks and breathing slowly through the nose for a time. When one is accustomed to keeping the



cheeks inflated, the mouthpiece of the blowpipe may be inserted between the lips, and the same thing repeated without attempting to blow or do more than keep the mouth full. Since the air now escapes through the blowpipe, the cheeks gradually fall together and must be again distended, yet without interrupting the outward current. This is accomplished by shutting off the communication between the mouth and the lungs by the palate, and inhaling through the nose. From the lungs thus filled the mouth is from time to time supplied, yet without any effort on the part of the muscles of the breast. A few hours' practice generally suffices to acquire the art of blowing. Beginners should keep in mind that the stream of air requires scarcely more force to produce it than results from the natural tandancy of the inflated cheeks to collapse. The lips should not be closed too firmly about the mouthpiece, else they are speedily fatigued. To the experienced operator continuous blowing is hardly an effort.*

THE FUEL.

9. When more convenient material is not at hand, stearine candles of good quality will answer for most purposes. Paraffine candles give a higher heat, but they soften in warm weather, and melt, and run down inconveniently. The common tallow candle may often suffice in an emergency, but requires constant snuffing.



10. A better fuel is olive or rape-seed oil burned in a lamp having a single circular wick rather more than a quarter of an inch in diameter, if the wick tube and lamp be so arranged that the charcoal and other supports used in blowpipe experiments can be brought close under the deflected flame. Fig. 5 represents the form of the blowpipe lamp proposed by Berzelius, and improved by Plattner. It is adapted for a portable blowpipe apparatus, since it is free from leakage, and capable of packing into a small space. The cistern A is of tinned sheet iron, and the wick tube and filling orifice are closed by screw caps.

11. The most convenient combustible is illuminating gas. A burner of the form given in Fig. 6 is used. It is about four inches high; the tube is flattened at the top and made a trifle lower on the left side, so that the blowpipe flame may be turned downward when necessary. A cock in the tube at the foot is useful. Such a lamp has the advantage of dispensing with all trimming and other inconveniences attendant on the use of an oil lamp. The ordinary Bunsen gas-burner (Fig. 7) is often provided with an extra tube to slip over the small gas jet in the in-

^{*} Luca has described a blowpipe intended to maintain a steady stream of air with intermittent blowing, but this and other contrivances are unnecessary when the student has sufficient enterprise and patience to learn to blow the ordinary instruments, and no others wilbe likely to make much progress in blowpipe analysis.

terior of the burner, in such a manner as to shut off the access of the air; the gas is then burned from the upper end, which is shaped as in the figure. The only objection to this lamp is, it is a little too high, although it may answer for all ordinary purposes.

A simpler blowpipe gas lamp may be easily made by selecting an iron or brass tube, eight inches in length, and three-eighths of an inch in bore, bending it at a right angle at the middle, and passing it through a block, properly cut, or placing it in a mould, which is then filled with melted lead. The top of the tube is then flattened, and the proper inclination given to the orifice by filing. Fig. 8 shows a lamp thus constructed.

THE BLOWPIPE FLAME.

12. When an ordinary lamp or candle is lighted, the combustion takes place only upon the outer limits of the flame, but if a stream of air is blown into the

flame the combustion is transferred to the interior, is thus rendered more complete, and the flame is condensed. It is to these causes that the very intense heat of the blowpipe flame is due.

When the beginner is able to maintain a steady blast for some minutes together, he may attempt the production and management of the blowpipe flame. The operator being easily seated at the table, his arm resting upon its edge, the blowpipe is lightly grasped near the water chamber, between the thumb and first and second fingers of the right hand, and its jet brought to the edge of the flame, just above the wick or tube. The blowing should be regulated so as to produce a steady flame, which will be regular and conical if the jet be well shaped.

In Fig. 9 a common candle flame is represented, in which a light-blue segment, bounded by the line a c, and disappearing as the



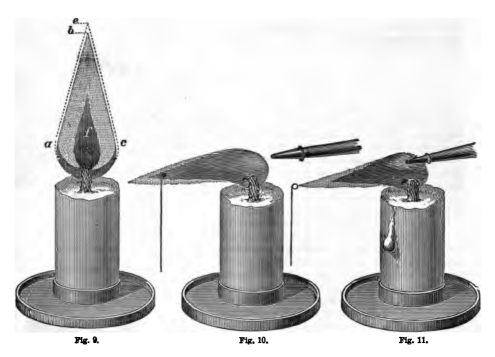
Fig. 8. (14 size.)

flame ascends, is seen at the base. The dark core of the flame f is surrounded by the illuminating portion a b c, and the thin, scarcely visible envelope a c c forms the outer coat of the flame.

13. Reducing Flame. While the candle is burning the stearine is slowly melted, sucked up by the wick, and vaporized. These vapors unite with the oxygen of the air and burn, upon the outer limits of the flame forming the hot coat a e c of carbonic acid and vapor of water. As the oxygen reaches no farther into the flame than the line a b c, the vapors inside this line are intensely heated out of the contact of the air, and any metallic oxide introduced into this yellow segment will, when hot, tend to part with its oxygen to the carbon and hydrocarbons of the flame. This is called the *Reducing Flame* (R. F.). To produce it with the blowpipe, the whole of the flame is deflected by a gentle blast, so regulated that it maintains its yellow color, and is luminous as before. The blowpipe is placed outside the flame, as shown in Fig. 10. The flame must not deposit soot upon the substance under trial, and only the extremity of the luminous part should be applied to as to envelop the assay.

SUPPORTS.

14. Oxidizing Flame (Fig. 11). When the jet is carried somewhat into the flame, and the blast is a little stronger, the carbon is more completely consumed; the inner blue cone, corresponding to the part a c of the candle flame, becomes sharply defined, and is surrounded by a nearly colorless envelope, corresponding to the mantle a c c, at the extremity of which metals may be intensely



heated in contact with air, and will thus be rapidly oxidized. This is called the *Oxidizing Flame* (O. F.). The assay should be held as far beyond the blue point of the flame as is consistent with the temperature requisite for the most rapid oxidation, and the flame should be so managed that no luminous streaks are seen in it. A flat wick serves best for its production. The heat is most intense at the point of the blue cone, and this is accordingly used for testing the fusibility of minerals and other substances, without reference to chemical action.

SUPPORTS.

In blowpipe examinations the assay is supported by certain substances which are either infusible, or have the power of sustaining a high heat without changing their form.

15. Charcoal is used in many operations as a support for the assay. For most purposes any piece of well-burned charcoal that does not snap or become fissured in the flame will suffice. The softer kinds of wood yield the most suitable material. That made from bass-wood (linden) is the best; pine and willow charcoal are also excellent. For use it is conveniently sawn into parallelopipedons,

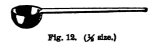
APPARATUS.

with faces one or two inches in width, and three to six inches in length. The assay is best placed on the flat, smooth surface, at right angles to the rings of growth. It can be repeatedly used, the clean surface being renewed by scraping with a knife or file.

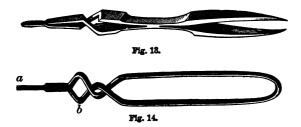
16. Cavities for the reception of the substance to be heated on charcoal may be made with the point of a knife. For some purposes, cavities may be made more nicely by means of a tube of stout tin plate, the edges of which are sharpened. The tube is made conical, has a length of three inches, its diameter at one end is three-eighths, at the other five-eighths, of an inch. The end of this is applied to the surface of the coal at a considerable inclination, and the tube is revolved with a scooping motion. The excavation should be made near the edge of the charcoal, should be cup-shaped, rather shallow, quite smooth, and regular.

17. Platinum Wire is used for supporting beads of fused borax or other flux in the flame. The kind designated as No. 27 (or jeweller's hole $12\frac{1}{2}$), is the best. It is cut into pieces three inches long, and a loop made in the end. When not in use the hooked ends should be plunged into a little bottle containing dilute sulphuric acid, which dissolves away the matters that have been fused on them. Before use they should be rinsed with water and thoroughly cleaned.

18. Platinum Spoon. For a few operations a small platinum spoon of the form shown in Fig. 12, may be usefully employed. A cork or wooden handle should be adapted to it. A rectangular slip of platinum foil, which is used also for other purposes, may be made to answer for the spoon by bending up its corners and holding it in the platinum forceps.



19. Platinum Forceps. For igniting fragments of minerals, forceps tipped with platinum are indispensable. Fig. 13 represents the usual form. They are made of steel or German silver. The points are opened by pressure. The free ends may be used as an ordinary forceps for picking up small fragments of minerals, etc.; or if of steel, for detaching pieces of specimens. Fig. 14 shows a



simpler form of this instrument, which any jeweller can easily construct. A piece of highly elastic brass wire, No. 12, is the best material for the tongs. The platinum tips a are readily hammered out from a piece of stout wire or cut from a plate, and are riveted or, better, soldered to the brass wire with silver solder. The bend at b is intended to prevent the points from touching the table. The forceps must be slender in order not to conduct away too much heat from the assay.

20. Glass Tubes. Tubes of hard glass, free from lead, $\frac{1}{18}$ to $\frac{1}{2}$ inch in diameter, and four to six inches long, are indispensable. They serve for the ignition

APPARATUS.

of bodies in a current of air, the rapidity of which may be regulated by varying the inclination of the tube. The substance under trial is placed in the tube about



an inch from the end, the tube is then held nearly horizontally, either in the flame of the lamp or of the blowpipe. The falling out of the body may be hindered by bending the tube slightly one inch from one end. The body is then placed at the bend as shown in Fig. 15, and the proper inclination given to the tube; but for most uses straight tubes are quite as good. For each new operation a clean tube must be employed. The tube usually cracks when used a second time, and should therefore be cut off at the place where a body has been ignited. Tubes are most easily cleaned by wiping them out with a slip of soft paper rolled around some slender cylinder having a *rough* surface to hold the paper. A small rat-tail file is excellent for this use.

21. Closed Tubes and Glass Bulb Tubes (matrasses; Fig. 16) serve for heating bodies out of contact, or with but limited access of air. They are easily made, especially the form B, which answers nearly every purpose, from the pieces which have become too short to be used as open tubes, or by heating a tube six inches long in the middle and drawing it into two parts.

ACCESSORY APPARATUS.

22. An Agate Mortar with pestle (Fig. 17) is used for reducing minerals to a fine powder. It should be from two to three inches in diameter, and should be used only for grinding, never for pounding, hard bodies.



23. A Diamond Mortar (Fig. 18) made of cast steel and well tempered, is used for breaking up and reducing to a tolerably fine powder hard and refractory bodies. The fragments to be broken are placed in the bottom of the cavity; the closely fitting pestle is also placed in the hole, and is sharply struck with a small hammer. Minerals are thus prepared for finer pulverization in the agate mortar; but the same thing may be accomplished by wrapping the assay in several folds of paper, placing it upon an anvil and striking it.

BLOWPIPE REAGENTS.

24. Hammer. A small steel hammer such as is used by jewellers.

25. Anvil. A small parallelopipedon of hardened steel, or any convenient flat surface of steel.

26. Pliers. Cutting pliers (Fig. 19) are useful for detaching fragments from mineral specimens.

27. File. A small three-cornered file is used for cutting glass tubes. A notch is cut in one side of the tube, which is then half pulled, half broken in two.

28. Magnet. A common steel magnet, or a magnetized knife blade, serves to recognize magnetic bodies; a magnetic needle is sometimes useful for delicate determinations.

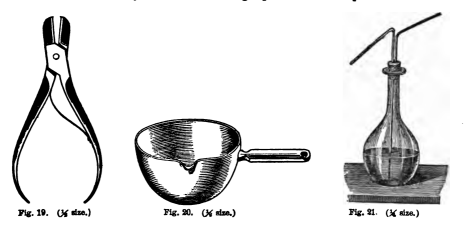
29. Lens. A magnifying glass composed of two convex lenses.

30. Watch-glasses from one to two and a half inches in diameter serve for various purposes.

31. Test-Tubes of hard glass with a suitable stand.

32. Funnels of glass one and a half to two inches in diameter.

33. Porcelain Dishes. Those with handles, called casseroles (Fig. 20), are most convenient. They are used for boiling liquids and for evaporations.



34. A Wash-Bottle (Fig. 21), made from a small flask, or any bottle having a mouth wide enough to receive the cork through which the tubes are passed.

35. Glass Rods, three to six inches long, rounded at each end, are used for stirrers.

36. Filters. Suitable paper is cut into circular pieces, the radius of which should be a half inch less than the side of the funnel in which it is to be placed. It is twice folded upon itself, thus forming a quadrant; this is opened so as to form a conical cup, having three thicknesses of paper on one of its sides, and one on the other. It is snugly inserted into a funnel, and moistened from the wash-bottle just previous to use.

The list of appliances for blowpipe analysis may be indefinitely increased, but the simplicity of a blowpipe outfit, in rendering it non-expensive and portable, is very desirable. A little ingenuity will supply the place of much apparatus.

BLOWPIPE REAGENTS.

The substances employed to produce chemical changes in bodies for their recognition are termed reagents. The quantities needed are so small that it is usually advisable to purchase most of them; but as it is often difficult to procure reagents of proper quality, simple directions for preparing some of them, and for testing their purity, are here given.

37. Carbonate of Soda; or Soda, in blowpipe language. Either neutral carbonate or bicarbonate may be used. To prepare it, take four or five ounces of commercial bicarbonate of soda, free from mechanical impurities, place it in a porcelain mortar, add a little distilled water, and pulverize finely. Bring it upon a large filter in a glass funnel, and allow the water to drain off. Successive additions of water, in quantities of about one ounce, are made, until a few drops of the drainings, caught in a clean test-tube, and acidified with nitric acid, give no precipitate, nor even the faintest turbidity with a drop of clear solution of chloride of barium. The washing often requires several days, and is sometimes not complete before half of the salt has been washed away. It is thus freed from sulphuric acid, which contaminates the commercial salt. Soda that is purchased as pure should be tested for sulphur and sulphuric acid, as described in 145, before trusting its purity. The salt as thus prepared is spread out upon paper and allowed to dry. Part of it may be bottled while moist, and used in that state ; but a part must be dried at a high heat, in order to expel all water. It is then pulverized and put away for use.

38. Biborate of Soda. Borax. The commercial salt is usually pure enough. Clean crystals are selected, and coarsely pulverized. For some tests, fused borax is required. To obtain this, some of the commercial salt is melted in a platinum dish, and when cool placed in a tightly stoppered bottle.

39. Phosphate of Soda and Ammonia. Salt of Phosphorus. Microcosmic Salt. The very small quantity of this substance (1 oz.) needed for a great number of trials is best purchased. It may be prepared by dissolving in two parts of boiling water six parts of crystallized phosphate of soda, and one part (all the parts by weight) of white and clean sal-ammoniac, and immediately filtering while still boiling hot. The crystals that separate on cooling are freed from the chloride of sodium that adheres to them by recrystallization. Testing.— It must fuse on platinum wire to a colorless, perfectly transparent globule; and when oxide of copper is added, and it is again heated, it must not tinge the flame with a blue or green color.

40. Nitrate of Cobalt. Cobalt Solution. The crystals of nitrate are dissolved in ten parts of water, and filtered if necessary. For use the cobalt solution

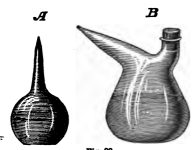


Fig. 22.

is most conveniently kept in bulbs similar to those represented in half size, in Fig. 22. The bulb A is easily made from a bit of glass tube. In order to fill such a bulb it is ge_tly heated, and the tip placed beneath the surface of a solution of nitrate of cobalt in a shallow dish. When a drop of the solution has entered it is again heated, the drop is converted into steam, the tip is again immorsed, and the solution will almost instantly rush into the bulb. It should not be more than two-thirds filled. To apply the solution, the bulb is grasped gently in the palm of the hand, and inverted, when

the expansion of the air shortly forces out a drop or more, as is required.
41. Nitrate of Potash. Clean crystals of the commercial salt are powdered.
42. Bisulphate of Potash. Equal weights of clean nitrate of potash and oil of vitriol are heated together in a porcelain dish, gently at first, afterwards more

strongly, till the nitric acid and excess of sulphuric acid are driven off and a clear liquid remains which solidifies to an opaque mass on cooling. The salt thus obtained is pulverized, and preserved in a glass-stoppered bottle. It can also be prepared by heating pure sulphate of potash with an excess of sulphuric acid, until

the excess is driven away and the mass solidifies on cooling.

43. Cyanide of Potassium. In nearly every case this reagent can be dispensed with, by one who has perfect command of the blowpipe, its only use being to facilitate difficult reductions. It can be procured of any photographer or druggist.

44. Iodide of Potassium. The clean crystals purchased of any druggist.

45. Sulphur. Flowers of sulphur.

46. Tin. Strips of pure tin-foil a half an inch wide and one inch long.

47. Zinc. Strips of common sheet zinc.

48. Lead. Pure lead, for detecting gold and silver by cupellation, is prepared by dissolving acetate of lead (sugar of lead) in hot water, filtering, and inserting strips of clean zinc into the solution. After five to six hours the precipitated lead should be scraped from the zinc in order to expose a fresh surface. When the lead is all separated, it is washed thoroughly with water, then dried by pressing between folds of blotting paper, and finally by exposure to a gentle warmth.

49. Iron. Clean wire of the thickness of a medium-sized sewing-needle. Iron in a fine state of subdivision is used for reductions in the wet way.

50. Magnesium. Bits of foil or wire are useful in detecting phosphoric acid. 51. Silver. A smooth silver coin, which must be freshly cleaned at the time of using. See detection of sulphur, 145.

52. Bone-Ash. A little cup of bone ashes, called a *cupel*, is used for the detection of silver and gold. Bones burned to whiteness are finely pulverized and reserved for these purposes.

53. Oxide of Copper. A copper cent is dissolved in nitric acid, the solution is evaporated to dryness, and the dry mass gradually heated to redness in a porcelain dish.

54. Fluoride of Calcium. Pure *fluor-spar* is crushed and heated in a test tube until decrepitation ceases; it is then finely pulverized.

55. Oxalate of Nickel. The pure salt is best purchased, and when fused with borax before the blowpipe, must give a brown and not a blue glass.

56. Test Papers. A. Blue Litmus Papers.—Digest one part of the litmus of commerce with six parts of water, and filter the solution; divide the intensely blue-filtrate into two equal parts; saturate the free alkali in the one part by repeatedly stirring with a glass rod dipped in very dilute sulphuric acid, until the color of the fluid just appears red; add now the other part of the blue filtrate, pour the whole fluid into a dish, and draw strips of fine unsized paper through it; suspend these strips over threads, and leave them to dry. When dry, the paper should have a fine blue color, and may be cut in narrow strips and kept in a tight box. The moistened paper should be promptly reddened by the smallest trace of acids, and is used for their detection. When the litmus paper is reddened by a very feeble acid, it may be used for the detection of alkalies.

B. Turmeric Paper.—Digest one part of bruised turmeric root with six parts of weak spirits of wine, filter the tincture obtained, and steep slips of fine paper in the filtrate. The dried slips must exhibit a fine yellow tint. It is turned brown by alkalies, and serves also in the recognition of boric acid, molybdic acid, and zirconia.

C. Brazil-wood Paper.—Brazil-wood is boiled with water, the liquid filtered, and paper saturated with it and dried. It is used for detecting fluorine, which gives it a yellow color; it also serves to recognize alkalies, which color it violet.

WET REAGENTS.

57. Water. Whenever water is used in analytical operations it should be

either distilled water, or clean rain water. 58. Hydrochloric Acid. Muriatic Acid. The strong commercial acid will answer for most purposes, but it is also advisable to have some of the pure fuming acid which on evaporation leaves no residue and when diluted with water gives no milkiness on the addition of chloride of barium.

59. Sulphuric Acid concentrated, (ordinary oil of vitriol).

60. Nitric Acid, pure. It must leave no residue upon evaporation, nor give any turbidity with nitrate of silver.

61. Phosphoric Acid. The officinal concentrated solution.

62. Ammonia. It must be colorless, should leave no residue when evaporated on a watch-glass, nor should it cause the slightest turbidity in lime-water.

63. Carbonate of Ammonia. One part of the commercial salt is dissolved in four parts of water, to which one part of solution of caustic ammonia has been added.

64. Chloride of Ammonium. Select sublimed white sal-ammoniac of commerce. If it contains iron or other impurities it is dissolved in hot water, and set aside to recrystallize. The dried crystals are dissolved for use in eight parts of water.

65. Phosphate of Soda. Purify the salt of commerce by recrystallization, and dissolve one part of the pure salt in ten parts of water.

66. Oxalate of Ammonia. Dissolve commercial oxalic acid, which has been purified by recrystallization, in two parts of hot water; add caustic ammonia, or carbonate of ammonia, until the fluid begins to manifest a slight alkaline reaction; filter, and set aside to cool. The crystals that separate are allowed to drain, and the mother liquors are further evaporated to crystallization. Purify by recrystallization. Dissolve one part of the pure salt in twenty-four parts of water.

67. Potassa. Dissolve some sticks of caustic potassa in water, allow to stand, and separate the clear solution from the sediment by decantation.

68. Chloride of Barium. This salt may be purchased of any druggist. For use it is dissolved in ten parts of water.

69. Nitrate of Silver. May be procured in crystals from any druggist or photographer.

70. Bichloride of Platinum. Treat platinum filings (purified by boiling with nitric acid) with concentrated hydrochloric acid and some nitric acid, and apply a very gentle heat, adding occasionally fresh portions of nitric acid, until the platinum is completely dissolved. Evaporate the solution to dryness on a water bath, with addition of hydrochloric acid, and dissolve the residue in ten parts of water for use. It is used for detecting potassa in the presence of soda and lithia.

71. Molybdate of Ammonia. Pulverize the sulphide of molybdenum as finely as possible, and roast it in a shallow sheet-iron or earthen dish, at a low red heat, until it turns yellow, and becomes converted into molybdic acid. It is then digested with ammonia, which extracts the molybdic acid; the solution is filtered, evaporated to dryness, and the molybdate of ammonia which is left is dissolved in water acidulated with nitric acid and kept for use.

All the reagents of a well-appointed laboratory may be of occasional service in the qualitative analysis of minerals, but reagents other than the above will be but rarely needed by the student in blowpipe analysis.

Chapter 2.

SYSTEMATIC COURSE OF QUALITATIVE BLOWPIPE ANALYSIS.

72. THE student being provided with the necessary materials, and having acquired some skill in producing the oxidizing and reducing flames, is prepared to consider the various effects that may be produced with the blow-pipe. These reactions are classified, according to the apparatus and reagents that are used, under the eight following heads, as recommended by Plattner:

A.--Heating in the closed tube.

B.—Heating in the open tube.

C.—Heating on charcoal.

D.—Heating in the platinum forceps to test fusibility, and to observe the coloration of the flame.

E.-Treatment with cobalt solution.

F.—Fusion with borax.

G.—Fusion with salt of phosphorus.

H.—Treatment with carbonate of soda.

Under each of the above divisions is given, first, the method of experimenting, and, second, in tabular arrangement, the phenomena or reactions produced, which are characteristic of the substances usually subjected to blowpipe examination. The beginner should not attempt at first to work with bodies of unknown composition, but should provide himself with some substances which are well calculated to illustrate the reactions indicated.

The blowpipe lamp is placed upon a sheet of stout clean paper, so that the assay accidentally falling may not be lost. Whenever a new substance is taken for experiment, all fragments of the old should be shaken off.

The assay must not be too large; in most cases the bulk of a mustard seed is enough, in the practised hand. Beginners may use a larger quantity, but as the student progresses he should aim to reduce the size of his assays to the least amount consistent with a perfect experiment, since he will be often called upon to determine minerals upon minute fragments.

The closest observation will often be necessary for the detection of the reaction, and the success of the student is greatly dependent upon the accuracy of discrimination, quick comprehension, and careful manipulation which is acquired in these preliminary examinations.

A.—HEATING IN THE CLOSED TUBE.

73. The body, in fragments the size of a grain of wheat, or an equivalent bulk of it, if it be in form of a powder, is placed in the bottom of a tube closed at one end; the tube is held nearly horizontal, and heated over the spirit or gas lamp,

very gently at first, and finally, if needful to intense ignition, with the aid of the blowpipe, and the successive phenomena are carefully watched as they appear.

Powdered substances must be so introduced into the tube as not to soil its sides; this is accomplished by placing the powder on a narrow slip of writing paper previously folded lengthwise in the form of a trough. The tube is held horizontal, and the paper trough is inserted to its bottom; the whole is now brought into a vertical position, and the paper is carefully withdrawn.

The phenomena can nearly all be produced in the simple closed tube (Fig. 16, b) and for most purposes this form is better than the bulb tube, since the object of these experiments is to heat the body out of contact with the air, and to produce changes among its constituents without the interposition of any reagents.

The following phenomena may be observed :*

1. Decrepitation—Fluorite, Barite, and many other minerals.

2. Glowing-Gadolinite, etc.

Phosphorescence—Fluorite, Willemite, etc.
 Change of color. The most important are here tabulated.

ORIGINAL COLOR.	COLO R While Ignited.	COLOR AFTER COOLING.	SUBSTANCE.
White to yellow.	Brown.	Yellow.	Binoxide of tin.
White.	Yellow.	White.	Oxide of zinc and many of ite
White.	Yellow.	Yellow.	salts. Hydrated oxide, carbonate,
Blue or green.	Black.	Black.	and other salts of lead. Hydrated oxide, carbonate,
White.	Dark yellow.	Light yellow.	and other salts of copper. Hydrated oxide, carbonate,
White.	Brown	Brown.	and many salts of bismuth. Hydrated oxide, carbonate,
Yellow or red.	Deeper color.	Original color if gently heated; green if strongly heated.	and many salts of cadmium. Most chromates.
Red.	Black.	Red.	Sesquioxide of iron.

5. Fusion-Stibnite, Nitre, and other bodies.

6. Give off oxygen-Binoxide of Manganese, Oxide of Mercury, etc. Tested by placing a bit of charcoal in the tube, upon the assay. Heat the charcoal first, then the assay, and the charcoal will glow.

7. Become carbonized, and give a burnt odor-Amber and many organic compounds. If acid reaction, non-nitrogenous; if alkaline, nitrogenous body.

8. Give off water—All hydrates.

9. Give acid vapors-Hydrates with volatile acids. Tested by placing a blue litmus paper in the end of the tube. If the glass is etched, Fluorine.

10. Give alkaline vapors—Ammonia Salts. Tested with a piece of turmeric paper.

11. Give sublimates which condense on the cold part of the tube.

* For experiments illustrating the effects to be produced in the closed tube, the following substances are given : Fluorite, Gadolinite, Oxide of Zinc, Stibnite, Oxide of Mercury, Amber, Serpentine, Nitrate of Ammonia, Pyrite, Bealgar, Arsenopyrite, Selenium, Amalgam, **Oinnabar**, Spathic Iron.

HEATING IN THE CLOSED TUBE.

a. A yellow sublimate.	} Su	LPHUR.	Either originally free, or from decomposition of a sulphide.
b. A sublimate, dark brown- red, almost black when hot, and red or reddish yellow when cold.	}	OF SENIC.	Realgar and Orpiment, and other Sulph-arsenides.
c. In strong heat, a sublimate deposits near the assay, which is black when hot, and brown- red when cold.		SULPHIDE OF CIMONY.	Sulphide of Antimony and its compounds, with other metallic sulphides.
d. A dark red, almost black, sublimate, and odor of decaying horse-radish at open end of tube.	SEL	ENIUM.	Various <i>Selenides</i> .
e. Condenses in small drops, with metallic lustre.	} Tell	LURIUM.	Various Tellurides.
f. A black, brilliant subli- mate, and garlic odor.	} Ar	SENIC.	Native Arsenic and many Arsenides.
g. A gray sublimate, con- misting (use lens) of metallic globules, which may be united by rubbing with a feather.	Mπ	RCURY.	Amalgam s.
h. A black, lustreless sublimate, red when rubbed.	}	LPHIDE OF RCURY.	Cinnabar, Vermilion, minerals containing both Mercury and Sulphur.
i. The body <i>fuses</i> , and yields a sublimate, which is white when cold.	$\begin{array}{c c} 2. & C \\ partiall \\ white o \\ 3. & A \end{array}$	y sublimes, n cooling. <i>ntimonous A</i> air be exclu	Mercury. ad; fuses to a yellow liquid, and becomes opaque and cid; fuses to yellow drops, uded, deposits in brilliant
j. The body does not fuse, but gives a sublimate, which is white when cold.	$\left.\begin{array}{c} 2. \ A\\ \text{denses}\\ 3. \ P\end{array}\right.$	in octahedral	onia. l; easily sublimes and con- crystals (lens). of Mercury; sublimate is

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12. The residue is magnetic—Spathic Iron, Pyrites, etc. There are some other reactions more rarely observed, particularly in the study of minerals. Osmic acid forms a sublimate of white drops, which possess a disa-greeable odor. Cyanogen, when liberated, is recognized by its peculiar odor. Iodine volatilizes in beautiful violet fumes. From some of its alloys Cadmium

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• .

volatilizes and condenses as a black metallic sublimate. Sulphurous acid is given off by sulphides in an amount proportionate to the oxygen which surrounds the assav, but the place for its observation is in the open tube.

B.—HEATING IN THE OPEN TUBE.

74. This is essentially a roasting or oxidizing process. The substance is placed in a glass tube open at both ends, at a distance of about one inch from the end, at which point a bend is sometimes made (see Fig. 15); but for most operations a straight tube is preferable. The heat should be gentle at first and only gradually raised, otherwise some bodies may volatilize without oxidizing, and give the same sublimate as when heated in a closed tube. By changing the inclination of the tube, the current of air through it may be increased or diminished, and the oxidation made to proceed more or less rapidly. Not too much of the substance must be taken, and if satisfactory reactions be not obtained from a fragment, it should be pulverized. Bodies which decrepitate and lose volatile ingredients by heating in a closed tube, must be finely pulverized at the outset, and introduced into the tube by means of a paper trough.

A slip of moistened litmus paper should always be placed in the upper end of the tube when experimenting on an unknown substance, and when vapors begin to arise, attention should be given to their odor, and to the sublimates which con-dense on the inner surface of the tube. Many of the phenomena encountered in this trial are identical with those obtained in the closed tube. Only such as are peculiar or characteristic are here noticed.*

1. Odors.-a. Sulphur and sulphides in the open tube form sulphurous acid, giving the odor of burning sulphur, and reddening moistened blue litmus paper. When a reaction is not developed by heating a fragment, the powder must be employed. b. Odor of decaying horse-radish.—Selenium. Mostly sublimes.

c. Odor of garlic.—Arsenic. Mostly sublimes.

2. Sublimates.—Carefully compare last section in case they are not noticed below. The sublimate itself should be heated to ascertain if it be volatile or fusible.

a. White, crystalline (octahedral), volatile ARSENOUS and many Arsenic and many Arsenic Acid.
b. White, thin sublimate, crystalline nearest the assay; fusible to droplets; yellowish when hot, nearly colorless when cold. When the R. F. is directed upon it within the tube, it be- comes blue, or even copper-red from reduction. MOLYBDIC Acidand Sulphideof Molybdenum.
c. Dense white smoke, and at first a mostly volatile white sublimate, depositing on the upper side of the tube; afterward in most cases a white, non-volatile and infusible sublimate gathering on the under side of the tube. ANTIMONOUS ACIDS. ANTIMONOUS ACIDS. Most compounds of Antimony.
d. White smoke, and non-volatile fusible SULPHATE OF Sulphide of sublimate depositing on the under side of tube. LEAD.

* Substances serving to illustrate the reactions of the open tube : Pyrite, Blende, Selenium Arsenopyrite, Molybdenite, Stibnite, Galenite, Bismuth, Tellurium, Cinnabar.

f. A gray sublimate, fusible to colorless drops that solidify on cooling. g. A steel-gray sublimate, the upper edge of which appears red, and sometimes fringed with SELENUE SELENUE SELENUE SELENUE SELENUE	npound s muth.
which approach and comparison fringed with Selenium	l many
small white very volatile crystals of selenous	
h. A bright metallic sublimate, that can be gathered into a drop by sweeping it together a splinter of wood or a feather.	ulphide

3. Residues.—Compare table of changes in color, 73. 4.

C.—HEATING ON CHARCOAL.

75. A small quantity of the substance is placed in a shallow cavity on charcoal, which is so situated that the flame of the blowpipe can be directed downward upon it, and its behavior in both flames observed.

A fragment may be used, or if the substance is in the form of powder, or on account of decrepitation must be reduced to powder, it may be mixed with water to a paste and placed on the coal, and heated at first gradually, afterward, when dry, to full ignition.

Much trouble is sometimes experienced in keeping the assay in its place sufficiently long to observe its behavior fully, especially when it is infusible or difficultly volatile. In such cases borax may often be employed to advantage in the following manner: The assay is held in the forceps, heated to redness, and then touched to a little grain of borax. The borax melts, and attaches itself to the body, which is now laid in the cavity so that the borax is in contact with the charcoal, and is carefully heated with the blowpipe; it usually adheres without further trouble.

In the following tables are given the characteristic phenomena that belong to this section.

1. Odors should be observed immediately after a short exposure to heat. Traces of sulphur, selenium, and arsenic are more surely detected by their odor on charcoal, than in an open tube.

- a. Odor of burning sulphur.-Sulphur and sulphides. Best observed in O. F.
- b. Odor of decaying horse-radish.—Selenium and selenides. Treat in O. F.
- c. Odor of garlic.—Arsenic and its compounds. Traces are most surely recognized after momentary exposure of the assay to the R.F.
- **2.** Deflagration.—Nitrates, chlorates, iodates, bromates.

3. The body fuses and is absorbed by the charcoal.—The fixed alkalies and many of their salts; also hydrates of baryta and strontia, and after very long heating their carbonates and sulphates.

- 4. A white infusible residue remains, it may be after previous fusion, which:
- a. Glows brightly in O. F., indicating lime, strontia, magnesia, zirconia, zinc, and tin.
- After ignition turns moist turmeric paper brown. Baryta, strontia, lime, magnesia.
- c. Communicates a characteristic color to the flame. See page 20.

	ИЕАВ АВВАТ.		DINTANT FROM ASSAT, OR DI THUN LAYERS.	IN O. F.	LAR.F.	REMAND.
a. Bet.swittet.	Bteel gray; fain	sel gray ; faint metallio lustre.	Dark gray, with tinge of violet;	Volatile	Volatile with blue flame.	Belenium fuses very easily ; volatilizes with brown- smoke, giving the odor of decaying home-radials.
A. THELORIDM.	White	•	Red or deep	Volatile.	Volatile with green fame, or, if selenium be present, with blue-green	Tellurium fuees very casily.
e. Aberrio.	White.		Graylah.	Volatile.	Volatile, with faint blue flame.	Metallic arrendo volatilizes without fuzing "ubli- mate is deposited quite from assay, is very vola-
4. ANTIKONT.	White.		Bluish.	Volatile.	Volatila, with faint greenish flame.	Luk and III. I. F. R. F. Rein gerund out. Metallio antimony fuses very early ; after being strongly basked nono charcoard, remains red-hot for a considerable time, and before solidifying becomes surrounded with crystals of antimonous scil. The
G. TEALLIUK,	White.		Near the assay	Volatile.	Volatile, with intense	Thallium fuses and oxidizes very cardiy.
/. Вщугв,	Reddiah brown; woken a kutle lead and antimony are present, carmine red. Eot. Cold.	when a little my are presend, Cold.				Bilver funes.
6. Bisnoria.	Dark orange yellow.	Lemon yellow.	Bluish white.	Volatile.	Volatile.	Blemuth fuses very easily. When subplate and cliorade of blemuth are submitted to the blowpipe on obsrooal they fuse, and outside of the sublimate of ordio is deposited a works consing of subplate or cliorade of blemuth, which is volatile in the B. P.
A. LEAD.	Dark lemon yellow.	Sulphur yellow.	Bluish white.	Volatile.	Volatile, with asure- blue flame.	wingurb counting Liead frame easily. When supplide and chloride of lead are basied B. B. on chronoad, they frame, and do- post a uplide sublimate of surplace or chloride of lead and or ording of the costing of ordios. The white sub- limited is sublide in P. B. thermothe dame hite sub-
6. Lindrak.	Dark yellow.	Yellowish white.			Volatile, with a violet	
J. CADAGUNE.		Bed brown.	Orange yellow.	Volatila.	name. Volatile.	Cadmium fuses easily, is volatile in R. F., and burna in O. F. with dark-yellow fame and brown smoke. The charcoal exterior to the sublimate some-
A, Zunc.	Yellow.	White.		Non-volatile, but glows brilliantly. After moisten- ing with nitrate of cobalt- and strong ignition be-	Blowly volatile.	and counter interscells. Chrynesty counter inter- section of the section of the sublimate of ordio s white constant of chirolice that is volatile in R. F. Zinc these easily, is volatile in R. F., and huran in O. F. with a imminus greenian-white finane. Chartes of fine finese, is partially decomposed, and partially ordiones unchanged in form of a white antiburge
1	Faint yellow.	White.		come success of each of the second se	ģ	Outdoor the controp of others. It is worked in it, a: This fitues easily, and in O. F. becomes covered with oride, which may be blown away mechanically. In B. F. the fured metal remains brilliant and the char- coal is conted. <i>Chiorids of its</i> behaves like chioride of the conted.
MOLTBURNE	Yellow ; some- times crystalline.	White.	Bluish.	Volatile, leaving a cop- leaved stain of oxide of molybdenum which is not further affected.	Gives a beautiful asure blue when touched for a moment with the B. F. : continued heat gives the same as in O. F.	Volatific, leaving a cop- Gives a beautiful arme of this sublimate is best obtained with Fulverised parted stain of oride of blue when touched for a material. When subhide of molybdenum is beated molybdenum which is moment with the R. F.; B. B. a copper-red ring surrounds the assay interior not further affected. [contured hest gives the to the white sublimate. Molybdenum is infusible.]

TABLE OF COATINGS GIVEN ON CHARCOAL

18

5. Sublimates or Coatings.—The volatile metals and some of their compounds give B. B. on charcoal, more or less characteristic deposits or sublimates. These coat the charcoal at a greater or less distance from the assay, and it must be observed what color they possess both when hot and cold, as well as whether they disappear in the O. F. and R. F., and thereby color the flame.

These sublimates, which are mostly deposited on the unheated charcoal, are not to be confounded with the ash (usually white), which remains as a thin coating where the coal itself has been exposed to the blowpipe flame.

Compounds of some of the metals must be heated in the R. F. They are then reduced to the metallic state, volatilized, and issuing from the flame are instantly reoxidized and deposited as a coating.

The characters given in the tables belong to the unmixed bodies. Their detection is often difficult when they occur together, and not always certain, even to the experienced operator.

a-m (inclusive). See table on page 18.

n. The sulphides (sulphates which in R. F. on charcoal become sulphides), chlorides, iodides, bromides of potassium, sodium, rubidium, and cæsium give B. B. white sublimates, the similar compounds of lithium grayish white, less copious sublimates, the salts themselves fusing and being absorbed by the charcoal. These sublimates volatilize in R. F., thereby tinging the flame with the color characteristic to these alkali metals: viz., potassium, rubidium, cæsium, violet; sodium, yellow; lithium, purple.

o. The chlorides of ammonium and antimony, and subchloride of mercury, volatilize without fusing, and yield white sublimates, which disappear in R. F. without coloring the flame.

p. Cheoride of copper fuses and tinges the flame intense azure blue. By long heating it partly volatilizes in white fumes, that smell of chlorine, and coat the charcoal with three differently colored sublimates, of which the interior is dark gray, the middle is dark yellow to brown, and the outer is bluish white. In R. F. the sublimate volatilizes, tinging the flame blue.

D.-HEATING IN THE PLATINUM-POINTED FORCEPS.

Coloration of the Flame.

76. Several bodies may be recognized by the colors they communicate to the blowpipe flame.

When the substance admits, a thin fragment may be held in the *clean* platinum forceps, and its point brought into the edge of the blue flame just within its apex.

When the body fuses so readily that it cannot be supported in the forceps, or if it attacks platinum, it must be laid in a very shallow cavity made on a narrow piece of charcoal, and held in such a manner that the flame may be thrown across it.

If the assay is infusible and decrepitates, or cannot be had in fragments, its powder is moistened to a paste with pure water (not with saliva) and spread upon the coal; it is first dried by a gentle heat and afterwards strongly ignited. Usually a coherent cake is thus obtained, which, with care, may be lifted in the forceps and its edge subjected to the flame. If a small fragment of a decrepitating mineral is taken in the forceps, and the forceps inserted into the flame in such a manner as to strongly heat their points before the mineral is heated, it may then be slowly drawn into the flame, uniformly heated, and thus often be saved.

The trial often succeeds best when the loop of a platinum wire is moistened with distilled water, touched to the powder of the assay, and then carefully heated; or if the body is easily fusible the wire may be ignited and brought rapidly in contact with it. Enough will adhere to observe if any coloration be given. Even if the substance attacks or alloys with the platinum, this method is to be recommended; it is then only needful to cut off the injured part of the wire.

The utmost care must be taken that no foreign matters interfere with the observation. The forceps, charcoal, or wire must be chemically clean, and must not alter the color of the flame when heated alone therein. If the assay is to be pulverized, the mortar and pestle must be thoroughly washed beforehand. The wire may be cleaned by dipping it in hydrochloric acid, or heating it therewith in a test tube, until apparently clean, and then rinsing it with distilled water. Merely by drawing a wire through the fingers, or wetting it with saliva, it receives a coating of soda enough to give a distinct though momentary yellow color to the blowpipe flame.

The flame itself should be what has been described as the oxidizing flame; it must at least be totally free from yellow streaks, and is best obtained from a slender wick like that of a candle. A brass wick-tube often tinges the flame green, especially if the fuel be oil.

The assay is held just within the point of the blue flame; the coloration is observed in the exterior part of the flame, and is best seen in a darkened room, or at least in a situation shielded from the direct light of day.

If the body gives no coloration or only a slight one when heated alone, it should be moistened with *sulphuric acid* and again heated, by which means phosphoric and boric acids become evident; or with hydrochloric acid, which in most cases heightens the coloration given by baryta, strontia, and copper.

1. YELLOW.

Reddish yellow. SODA in all its compounds, even when present in very small quantity. Admixtures of potash, etc., even in considerable quantities, do not in terfere with this reaction.

2. VIOLET.

Bluish violet. POTASH and most of its salts, phosphates, borates, and infusible silicates excepted. In presence of very little soda the reaction of both is discernible; with more soda (1 per cent.) the yellow flame predominates. The presence of lithia also masks this reaction. Silicates containing potash only, give the flame • violet color, when, besides being free from soda and lithia, they are somewhat fusible. Indium, cæsium and rubidium also give violet flames.

3. RED.

a. Purple red. LITHIA and most of its compounds. The reaction is not masked by potash, but easily by soda.

b. Red. STRONTIA and many of its compounds. The coloration is increased by moistening the already ignited assay with hydrochloric acid; is masked by much baryta.

c. Yellowish red. LIME and many compounds; flame not to be confounded with that produced by strontia; is masked by much baryta.

4. GREEN.

a. Yellowish green. BARYTA and most of its salts, silicates excepted; not masked by lime.

b. Yellowish green. MOLYBDIC ACID; also oxide and sulphide of molybdenum.

c. Emerald green. COPPER and most of its salts. d. Green. TELLUROUS ACID.

e. Green. THALLIUM and its salts.

f. Bluish green. PHOSPHORIC ACID. Many phosphates give the coloration alone; others only after their powder is moistened with sulphuric acid to a paste, and then ignited on platinum wire. The coloration is often but momentary. g. Yellowish (siskin) green. BORIC ACID. Minerals and salts are best mixed as

powder with sulphuric acid, and heated on platinum wire; coloration often momentary.

h. Dark green, feeble. Ammonia Salts.

i. Whitish green, intense. METALLIC ZINC.

5. BLUE.

a. Light blue. METALLIC ARSENIC, and arsenides of bases which do not them selves tinge the flame. Also arsenates, and arsenous acid.

b. Greenish blue. METALLIC ANTIMONY, and the sublimate of antimonous acid, on charcoal.

c. Azure blue. LEAD. The metal fused in R. F., the sublimate of oxide, also salts of lead when fused on wire, in case their acid constituent does not tinge the flame strongly.

d. Azure blue. SELENIUM. e. Azure blue. CHLORIDE OF COPPER. Metallic copper, and most copper compounds after wetting with hydrochloric acid, color the flame for a short time purplish blue, afterwards green.

f. Greenish blue. BROMIDE OF COPPER. After a little time, green.

Fusibility.

77. The fusibility of minerals is also tested in the platinum forceps. (See scale of fusibility in Chapter IV.) As a general rule, no substances with metallic lustre should be heated in the platinum forceps, since they are apt to be injured by forming an alloy with the fused metals; but the cautious manipulator may heat any substance in the forceps without danger, by taking especial care that the fused substance does not come in contact with the forceps.

78. Many of the combinations of the alkaline earths become alkaline on heating. Such substances, if not too fusible, may be treated in the forceps, and the fragment under examination after cooling placed on a strip of moistened turmeric paper, which acquires a brownish-red color at the point of contact with the assay.

E.—TREATMENT WITH COBALT SOLUTION.

79. This operation is only applicable to bodies which are nearly or quite infusible, and which, after ignition, have a white or at least a grayish color, and is always conducted in O. F. If the substance can be heated in the form of splinters or fragments, and is somewhat porous, it may be held in the platinum forceps; the projecting extremity is moistened with the cobalt solution, then heated gradually until dry, and finally ignited as strongly as possible in O. F. without causing fusion.

Hard, compact minerals must be finely pulverized before treatment. The powder is placed in the palm of the hand and moistened with the solution of cobalt. A portion of the paste is then taken upon the loop of a platinum wire and strongly ignited in the O. F.

Certain sublimates, for example, oxides of zinc and tin, formed by heating compounds of these metals on charcoal, are treated directly with cobalt solution.

By this treatment several bodies, especially alumina, magnesia, and oxide of zinc, assume characteristic colors. The tints of blue, red, and black that appear before strong ignition are merely due to the drying or decomposition of the nitrate of cobalt, and are not to be regarded.

The color of the assay thus treated must be examined by daylight.

Minerals, and salts which fuse to a colorless glass, yield with cobalt solution the smalt-blue color which is characteristic of cobalt. A blue *infusible* mass only, indicates *alumina*.

The cobalt solution should be rather dilute, and if needful, successive portions added until decisive results are obtained.

This reagent serves to detect alumina, magnesia, etc., infallibly when they are in the pure state, and also in many of their combinations; but in various minerals the result is masked by other ingredients.

80. The colors thus obtained are given in the following table:

- 1. Brown or brick red-Baryta, under fusion and while hot.
- 2. Flesh red-Magnesia, tantalic acid, after cooling.
- 3. Violet-Zirconia (dirty violet); phosphate and arsenate of magnesia (fuse).
- 4. Blue-Alumina, silica (faint).
- 5. Green—Oxide of zinc (yellowish green), oxide of tin (bluish green), titanic acid (yellowish green), columbic acid (dirty green), antimonic acid (dirty dark green).
- 6. Gray-Strontia, lime, glucina (bluish gray).

It sometimes happens that the ash of the charcoal itself acquires a new color by ignition with this reagent. We have occasionally observed a greenish-yellow color thus produced. The operator has to assure himself that the ash of the coal he uses gives no deceptive reaction with nitrate of cobalt.

Use of Fluxes-Roasting.

81. Borax as well as salt of phosphorus exerts a very powerful solvent action when fused with metallic oxides, forming, in many cases, highly colored glasses, which are exceedingly characteristic. These salts are therefore very important reagents in blowpipe analysis; but it must always be remembered that the colors noted in the following tables are those given by the *oxides*, and where the preliminary examination has shown the substance to contain sulphur or arsenic in combination it is indispensable before going further to remove these elements, and convert the metals into oxides by *roasting*.

82. Roasting. The operation of roasting is performed as follows: The finely pulverized substance is placed in a quite shallow cavity on charcoal, pressed with a pestle or knife-blade into a *thin* layer, and heated for some time, only to dull redness, with the extreme point of the flame. When the odor of sulphurous acid ceases to be perceptible the assay is brought into the R. F., whereby the sulphates and arsenates that may have been formed in the O. F. are reduced,

and arsenic is more or less driven off. When no more arsenical odors are evolved the treatment in O. F. is repeated, and these operations are alternately continued until the assay is odorless in both flames. The heat should be quite moderate, so that the body does not fuse; if it fuses, it must be removed to the agate mortar and freshly pulverized. When the roasting has been well conducted the residue is pulverulent, and of uniform appearance throughout. When much arsenic is present it is best to heat the body previously in the open glass tube.

Bodies containing selenium, tellurium, and antimony, if free from sulphur and arsenic, usually require no roasting, as the former substances, unlike the latter, do not interfere with the reactions about to be described.

F.—FUSION WITH BORAX.

83. Treatment with Borax in O. F. The fusion with borax is usually effected on the platinum wire. The clean loop is heated to redness and dipped in borax powder, and the adhering particles are heated until fused to a clear and colorless glass, or bead; this bead, while still hot, is brought in contact with a very little of the assay, and heated therewith in the O. F.

It is to be observed whether the body dissolves readily or slowly, quietly or with effervescence; and when solution has been effected, the bead is to be held before the eye, against the light, and its color, when hot and cold, is to be noted, as well as whether its transparency is disturbed while cooling. Beads should not be looked at against the light of the gas or candle, since by such lights the colors are much modified.

The phenomena of color vary in intensity, and to a certain degree in kind, according to the quantity of substance dissolved in the bead. The manifestation of opacity on cooling depends also upon the quantity of material contained in the flux, and indeed only occurs when a certain amount has been added. It is therefore necessary to begin by dissolving a little of the assay, and after noting the result, more may be cautiously added at several intervals, until the operator is satisfied.

If, by using too much of the assay, a bead has been obtained, so deeply colored that it is difficult to decide what the color is, it may be flattened in the forceps, or drawn out by a platinum wire while still hot; or most of the hot bead may be thrown off with a sudden jerk, and the remaining portion diluted with more borax.

If the operator be in doubt as to the nature of the color he has obtained, he should view it through a lens, or compare it with some known color, obtained by fusing the appropriate pure metallic oxide in another borax bead. Care must be taken finally to guard against deception arising from reflections from colored surfaces near the operator.

84. Flaming. The alkaline earths, and some other bodies, dissolve in borax, forming beads which, at a certain stage of saturation, are clear, and remain so when cold, but which, if heated slowly and gently in the R. F., especially with an intermittent flame, become opaque and enamel-like.

The application of the *intermittent flame* is called *flaming*. In most cases the bodies, which at a certain degree of saturation are made opaque by flaming, become so without flaming when the saturation is carried a little farther.

85. Treatment with Borax in R. F. After observing the behavior of a body in the O. F., it is subjected to the R. F., which must, however, be so managed that no soot deposits on the bead. After blowing a little time the bead is allowed to cool, and its color, both when hot and cold, is observed. It may sometimes be needful to add more of the assay, and repeat the heating. In case no effect be pro-

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duced, or if metallic globules appear, which may often alloy with the platinum (whereby the loop is spoiled), the bead is jerked off into a clean dish, placed in a shallow cavity on charcoal, and further submitted to the R. F. for one or two minutes. In this way reductions are easily accomplished that scarcely succeed on the wire. While the bead is still glowing it is grasped in the clean pincers, flattened, and slightly lifted from the charcoal. It is thus suddenly cooled, whereby oxidation, that might occur were the bead left to cool slowly, is prevented, and at the same time it is brought into a good position for examining its color.

In special cases reduction is still further aided by help of metallic tin. A bit of tin-foil is laid in contact with the bead, and the two are fused together for a few moments in the R. F. The tin oxidizes at the expense of the higher oxide present, reducing the same to a lower oxide, while the oxide of tin formed, dissolves in the borax, without interfering with the color produced by the reduced assay.

86. With Borax in O. F. are yielded-

1. COLORLESS BEADS BY

TEMPERATURE.

Hot and Cold.	Silica, alumina, oxide of tin, baryta, strontia, lime, magnesia, glucina, yttria, zirconia, thoria, oxides of lanthanum and silver, tantalic, columbic, and tellurous acids:	when strongly saturated become opaque white by flaming.
Cola.	Titanic, tungstic, molybdic, and antimonic acids, oxides of indium, zinc, cadmium, lead, and bismuth:	when slightly saturated.

2. YELLOW BEADS BY

	Titanic, tungstic, and molybdic acids, oxides of zinc and cadmium :	flaming.
Hot.	Oxides of lead and bismuth, antimo- nous acid:	less when cold.
	Oxides of cerium, uranium, and iron :	when feebly saturated; paler on cooling.
	Oxide of chromium : when feebly satur	rated ; vellowish green when cold.

Oxide of chromium: when feebly saturated; yellowish green when cold. Vanadic acid: greenish when cold.

3. RED TO BROWN BEADS BY

		cerium:			

	" didymium; rose colored; the same when cold.
Hot.	" iron: yellow when cold.
not,	" uranium: yellow on cooling; opaque yellow by flaming.
	" chromium : yellowish green when cold.
	" iron containing manganese: yellowish red on cooling.
	Oxide of nickel (red brown to brown): violet when hot.
Cold.	" manganese : (violet red) violet when hot.
Cold.	" nickel containing cobalt: (with little cobalt, violet brown)
	violet when hot.
	-

FUSION WITH BORAX.

4. VIOLET (AMETHYSTINE) BEADS BY

(Oxide of nickel: red brown to brown on cooling.

- manganese: violet red on cooling.
 mickel containing cobalt: passes into brown on cooling; if much
 cobalt be present, it remains violet.
 cobalt containing manganese: on cooling, like the nickel mixture.

5. BLUE BEADS BY

Hot.-Oxide of cobalt: unchanged on cooling. Cold.-Oxide of copper (when highly saturated greenish blue): green when hot.

6. GREEN BEADS BY

(Oxide of copper: blue after cooling, or greenish blue when highly saturated. ... **.** . ..

Hot	Oxide o " "	of iron con " copper "	taining " " "	g cobalt: copper: iron: nickel:	According to the degree of saturation and the relative proportions of the oxides to each other, the green co- lor changes on cooling into pale green, blue, or yellow.
Cold.	∫Oxide (of chromiu	ım (ye	llowish gr	een): yellow to red when hot.
	Vanadi	ic acid (gro	eenish)): <i>yellow</i>	when hot.

87. With Borax in R. F. are given-

1. COLOBLESS BEADS BY

Hot	Silica, alumina, oxide of tin, baryta, strontia, lime, magnesia, glucina, yttria, zirconia, thoria, oxide of lanthanum, oxide of ceri- um, tantalic acid : Oxide of didymium, oxide of manganese, the latter often takes a faint
Cold.	rose color on cooling: Columbic acid: when used in small quantity.
	Oxides of silver, zinc, cadmium, lead, bis- muth, and nickel; antimonous and tellu- rous acids:
Hot.	Oxide of copper: becomes opaque red on cooling, if highly saturated.
	2. YELLOW TO BROWN BEADS BY

Titanic acid (yellow to brown) : when strongly saturated ; become enamel

blue by flaming. Tungstic acid (yellow to dark yellow) : brownish when cold. Molybdic acid (brown to black and opaque). Vanadic acid (brownish) : chrome green when cold. Hot.

3. BLUE BEAD BY

Hot. Oxide of cobalt : unchanged on cooling.

Hot.

FUSION WITH SALT OF PHOSPHORUS.

4. GREEN BEADS BY

	(Oxide of iron (yellowish or bottle green): especially when cold.
Hot	" uranium (yellowish green): when highly saturated; becomes
and	black by flaming.
Cold.	" chromium (pale to dark emerald green): according to degree
	of saturation.
Cold.	Vanadic acid (chrome green): brownish, when hot.

5. GRAY OF TURBID BEADS, THE TURBIDITY OFTEN APPEARING DURING THE HEAT-ING BY

Oxides of silver, zinc, cadmium, lead, bismuth, and nickel, antimonous and tellurous acids: Columbic acid: when highly saturated.

6. RED BEADS BY

Cold. { Oxide of copper (opaque) if highly saturated, or with tin on charcoal. Sesquioxide of didymium (rose color).

G.—FUSION WITH SALT OF PHOSPHORUS.

88. The general rules given for fusion with borax apply here.

The salt of phosphorus when first heated fuses in its crystal water, and is so fluid that it easily falls from the platinum loop. If, however, a small quantity be first fused upon the wire until it ceases boiling, then the additional quantity needed will adhere without difficulty. The bead is best placed *over* the blowpipe flame, as the ascending vapors that are driven from the salt buoy up the bead and keep it from falling.

In general the behavior of the various bodies is quite similar to that with borax ; there are, however, characteristic differences, as the table shows.

Salt of phosphorus is especially useful in the detection of *silica*. Most silicates, when added to a bead of it and heated, are decomposed. The bases dissolve in the flux without interfering with its transparency (unless the substance is in too large quantity), while the silica, being almost insoluble, floats as a translucent yet distinct cloud in the bead. It is best observed when the bead is hot. If the alkaline earths be present, the bead becomes opaque on cooling, but this does not interfere with the test. It must be borne in mind, however, that silica is soluble, though but slightly, in salt of phosphorus, and small quantities may, therefore, be easily overlooked. Also that some silicates, especially those of alumina and zirconia, are with difficulty decomposed by it.

When phosphate of soda and ammonia is subjected to the action of heat, the ammonia escapes with the water of crystallization, and the readily fusible metaphosphate of soda is left behind. This is a powerful solvent, and its action in quite analogous to that of biborate of soda.

39. With Salt of Phosphorus in O. F. are given-

Cold.

1. COLOBLESS BEADS BY

Hot and Cold.	Silica (very slightly soluble). Alumina, oxide of tin (difficultly soluble). Baryta, strontia, lime, magnesia, glucina, yttria, zirconia, thoria, ox- ide of lanthanum, tellurous acid: Tantalic, columbic, titanic, tungstic, and antimonous acids, oxides of zinc, cadmium, lead, and bismuth: Silica (very slightly soluble). when strongly saturated; be- come opaque white by flam- ing. when not too highly saturated; otherwise yellowish to yellow and colorless only after cooling.
	2. YELLOW BEADS BY
Hot. Cold.	Tantalic, columbic, titanic, tung- stic, and antimonic acids, oxides of lead, zinc, cadmium, and bismuth :when slightly saturated, but col- orless when cold.Oxide of silver (yellowish) : when cold, opalescent.Oxides of iron and cerium : when slightly saturated ; become colorless on cooling (strongly saturated are red when hot, and yellow when cold).Oxide of uranium : yellowish green when cold.Vanadic acid (dark yellow) : paler on cooling.Oxide of nickel : reddish when hot.
	3. RED BEADS BY
Hot.	Oxides of iron and cerium: when highly saturated; becomes yellow after cooling. Oxide of didymium: rose color when saturated. " nickel (reddish): yellow when cold. " chromium (reddish): emerald green when cold.
TT .	4. VIOLET (AMETHYSTINE) BEAD BY
Hot.	Oxide of manganese (brown violet): pale red violet when cold.

5. BLUE BEADS BY

Hot.	Oxide of cobalt: color unchanged on cooling.
Cold.	" copper (when strongly saturated greenish blue): green when hot.

6. GREEN BEADS BY

	Oxide of copper: blue when cold (when strongly saturated, greenish blue).
	Molybdic acid (yellowish green): paler on cooling
Hot.	Oxide of iron containing cobalt.According to the degree of saturation, and the relative proportions"""""""""""""""""""""""""""""""""""
Cold.	Oxide of uranium (yellowish green): yellow when hot. (chromium (emerald green): reddish when hot.

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90. With Salt of Phosphorus in R. F. are given-

1. COLORLESS BEADS BY

Hot and Cold.	Silica (very slightly soluble). Alumina and oxide of zinc (difficultly soluble). Baryta, strontia, lime, magnesia, glucina, yttria, zirconia, thoria, oxide of lantha- num: Oxides of cerium, didymium, and manganese. Oxides of silver, zinc, cadmium, indium, lead, bismuth, tantalic, antimonous, and tellu- rous acids: Oxide of nickel (especially on charcoal).
	2. Yellow to Red Brads by
Hot.	Oxide of iron (yellow to red): when cooling at first greenish, then reddish. Titanic acid (yellow): violet on cooling. Columbic acid (violet brown): particularly on charcoal. Vanadic acid (brownish): chrome green after cooling. Titanic acid containing iron. Yellow): when cold, brown (blood) Tungstic """" fred Columbic """ (brown red): dark yellow when cold.
	3. VIOLET (AMETHYSTINE) BEADS BY
Cold.	{ Columbic acid (when highly saturated) : faint dirty-blue when hot. { Titanic acid (even by moderate saturation) : yellow when hot.
	4. BLUE BEADS BY
Cold.	Oxide of cobalt: same when hot. Tungstic acid: brownish when hot. Columbic acid (when very strongly saturated): dirty blue when hot.
	5. GREEN BEADS BY
Cold.	Oxide of uranium : yellowish green when hot. Molybdic acid : dirty green when hot. Vanadic acid : brownish when hot. Oxide of chromium : reddish when hot.
	AY OR TURBID BEADS, THE TURBIDITY OFTEN APPEARING DURING THE HEATING BY
Cold.	Oxides of silver, zinc, cadmium, in- dium, lead, bismuth, and nickel, anti- monous and tellurous acids: Reaction best obtained of charcoal. After long blow ing become colorless.
	7. RED BEADS BY
Cold.	Oxide of copper (opaque) when strongly saturated, or by aid of tin on charcoal. Sesquioxide of Didymium (rose colored).

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H.-TREATMENT WITH SODA.

91. No attempt is here made to tabulate the phenomena that may arise in the treatment of bodies B. B. with carbonate of soda. These phenomena have either been described in the foregoing tables (sublimation), or are somewhat uncertain in their production, especially by the beginner (formation of glass with silicates), or, finally, are of a general nature (reduction of metallic oxides). We therefore translate substantially what Plattner has written under this head. According to the nature of the assay, it may either *fuse together with* or *dissolve in* soda, as when containing earths or fixed acid; or a *metallic reduction* may occur if the assay consist of reducible metallic oxides.

92. Fusibility with Soda (O. F.). A large number of bodies have the property to unite with soda at a high temperature, and to give partly fusible, partly infusible, compounds.

The fusible bodies are, however, few in number: principally silica, titanic acid, tungstic acid, and molybdic acid. When the fusion takes place on charcoal, silica and titanic acid both unite with the soda under effervescence to clear beads. The silicate of soda remains transparent after cooling if no excess of soda be present, but the titanate of soda becomes crystalline and opaque.

Molybdic and tungstic acids also combine with soda with effervescence, but the compounds are absorbed by the charcoal. Besides these acids, the salts of baryta and strontia are fusible with soda, but the mass is absorbed by the coal. Most salts of lime fuse indeed with soda, but when the acids they contain are stronger than carbonic acid, they are decomposed; the resulting salt of soda penetrates the coal, while the lime remains as a white mass on the surface.

In trying the fusibility of a body with soda, one proceeds in the following manner: If the body be in form of powder, it is mixed in the palm of the hand with soda, by means of a moistened knife-blade, to a coherent mass; if the assay be a splinter or fragment, and does not decrepitate, the moistened soda is spread upon it; if it decrepitates it must be pulverized. In both cases the assay is placed in a shallow cavity on charcoal, gently heated until thoroughly dry, and thereupon intensely ignited in the O. F. If a fragment has been used, the soda is commonly absorbed by the coal as it first fuses; but if the assay be soluble in it, it appears again and attacks the body with effervescence, and presently fuses with it to a globule. If too little soda be used in the treatment of a body soluble in this reagent, a portion of the assay remains undissolved, and surrounded by a clear glass; if too much soda has been employed, the glass will become opaque on cooling. It is therefore advisable to add the soda in successive small quantities, and observe the changes thus produced. Many bodies, especially silicates, which are themselves difficultly fusible, although their bases are infusible, dissolve in a little soda to a clear glass, but with more soda they form a slaggy or infusible mass.

If the assay be insoluble in soda, but decomposable by it, the operator will see that it gradually swells up and changes its appearance, though it does not fuse to a globule. If this be the case with an assay used in the state of powder, it may not be certain that it is actually insoluble, because too little soda may have been used; the mass must therefore be heated with a new portion of soda, or even with a second or third addition. When this appearance of decomposition occurs with a fragment of mineral, the same body must also be heated with soda in the state of powder. If the assay is both insoluble and undecomposable, the soda is absorbed by the charcoal and the body is left on the surface unchanged, whether applied as a fragment or in powder.

93. Formation of a Hepar (R. F.). The higher sulphides of the alkalies

have long been known by the name of *Hepar sulphuris* (liver of sulphur), since they possess a *liver-brown* color. When soda is fused on charcoal in the R. F. with any compound of sulphur (sulphide or sulphate), sulphide of sodium is produced, and if much sulphur was present in the assay the fused mass will show the characteristic color of hepar. Whether or not the mass possess this color, whether it remain on the surface of the coal or be absorbed by it, it is only necessary to place it on a freshly scraped surface of silver (or to cut out the coal into which it has sunk, and put it on the silver), and then add a drop of water, in order after a few moments to recognize the slightest trace of SULPHUR by the production of a yellow or even black stain of sulphide of silver. Illuminating gas commonly contains sulphur-compounds, and when this test for sulphur is employed with gas for fuel, the soda should always be fused first on coal and tested before adding the assay. If sulphur should prove to be present the test must be made with a candle or oilflame.

94. Reduction of Metallic Oxides (R.F.). The fusion of certain oxides with soda on charcoal in R.F. furnishes a most ready and delicate means of detecting their presence in minerals and salts.

Some metallic oxides are reduced to the metallic state by heating alone in R. F. when pure, but with difficulty or not at all when mixed or combined with other bodies; by addition of soda, however, the reduction is easy. There are other oxides that alone are unaltered, but by fusion with soda are reduced to the metallic state.

If the oxide of lead, for example, is fused with soda, there is no difficulty in recognizing the metallic lead, which will be found in globules on the surface of the charcoal. Oxide of iron yields, however, metallic iron which cannot be fused, and the fusible metals often escape the eye when present in small quantity. The operator must therefore employ the method of Gahn, as follows. The finely pulverized substance is mixed with soda and a drop of water to a paste, which is laid in a cavity on charcoal, and strongly heated in the R. F. The soda commonly sinks into the charcoal; more is added at intervals, until the assay has nearly or completely dis-appeared in the pores of the coal. A drop or two of water is now put upon the place, and all those parts of the coal near the cavity which have absorbed the assay are cut out into the agate mortar, and pulverized with addition of water to a fine powder. The water is now carefully decanted, or the mortar is held beneath the surface of water contained in a clean bowl, and gently moved to and fro, so that the coal dust is washed away from any metallic particles that may be in the mortar. By careful washing even the smallest quantity of copper, tin, or lead may be seen remaining in the mortar in the shape of flattened globules. If the metal be infusible or brittle, it will be found as a heavy, lustrous powder.

The nature of the metal can be determined by its physical properties; or the particles may be dissolved in borax or salt of phosphorus, and tested as already described. Often the sublimate that is deposited about the assay will give a clue to the kind of metal under examination.

Iron, cobalt, and nickel are obtained as metallic powder which is lifted by the magnet (best tried under water). Copper is recognized by its red color; Tin and Lead flatten under the pestle; Bismuth and Antimony are brittle, and present themselves as powder. Besides these metals, Molybdenum, Tungsten, Tellurium, Indium, Zinc, and Cadmium, and the noble metals, are also reduced by treatment with soda. Antimony, Tellurium, Bismuth, Indium, Lead, Zinc, and Cadmium volatilize partly or completely, and yield characteristic sublimates. Zinc and Cadmium usually volatilize entirely. Arsenic and Mercury are also reduced, but must be heated with soda in a tube, in order to collect the sublimates, which are metallic arsenic and mercury. When several metals are together, they usually form an alloy. Copper and iron are, however, obtained distinct. If the assay contained arsenate of cobalt or nickel, fusible metallic globules are obtained, which are always brittle from presence of arsenic. The reactions with borax and salt of phosphorus must be the final resort, and it may happen that only the experienced operator will be able to make out satisfactorily the nature of a metallic mixture, such as may result from a reduction with soda.

PLATTNER directs attention to the three following points, as needful to be carefully attended to in successfully conducting the operation in question:

1. The operator must keep the assay a sufficiently long time exposed to the action of a strong R. F.

2. In cutting out and pulverizing the fused mass, and in washing the same, the greatest care must be exercised that no metallic particles be lost; and,

3. The remaining metal, whether in form of scales, grains, or powder, must be examined with help of a lens, and tested by means of the magnet, and if needful by fluxes (borax and salt of phosphorus).

To acquire skill in the detection of copper and tin by reduction with soda (it is most applicable for finding small quantities of these metals especially), the beginner should practise with mixtures of a copper ore or salt with increasing quantities of feldspar or some other body free from metallic oxides. One or two per cent. of tin, and much less copper, can be detected in the quantity usually employed for blowpipe assays.

95. For convenience of reference is added here a tabular view, translated from Plattner, of the behavior of the earths and metallic oxides when treated successively, (1) alone on charcoal or in the platinum forceps; (2) with borax, and (3) with salt of phosphorus on platinum wire; (4) with soda, and (5) with cobalt solution; the special reactions of the alkalies will be given under a propriate heads in the next chapter. In the table the sign O, given under some of the heads, indicates that no reaction is observed with the substance.

Martha	Behavior alone, on Charcoel, and in the Flatinum forcepe.	With Borax on Platinum Wire.	With Salt of Phosphorus on Pla- thum Wire.	With Bods on Charcoal.	With Bolution of Cobalt in 0. 7.
Bacyta Ba	A hydrate fuses, bolk and intra- mesces, congreak on the surface, and then is absorbed by the onal. As carbonate, these cashy to a clear glass, which becomes trypeted fusion to colling. After repeated fusion it bolks and spirrs, becomes causide and is absorbed by the cost. Heated in the forceps, tinges the fame yellowich green.	As hydrate three, bolis and intri- meeces, congreak on the surface, and dhen is aborbed by the earbonate is soluble with at and dhen is aborbed by the coal. As earbonated in the earby and opeque by fam- bo a dear glass, which become trepeated fraton it bolis and spirity become caused and pellowish greet.		Tues together with sola, and is Fuses to a paie brown or brown a base to a paie brown or brown of giobule; on cooling long long long, and on exposure to the atmosphere falls into a light gray powder.	ruses to a pule brown or browniah rud giobule; on cooling losse coior, and on expecture to the simosphere falls into a light- gray powder.
	The hydrate behaves like that of As Baryta, beryta. The extromate these on coal only on the finest edge, and throws out cauliforwar-like ramifications which emits a bril- liant light, and they also react alla- haintly red; they also react alla- the with turner to paper. Heat- ed in the forenga, the fiame is tinged purple red.	As Baryta.	As Baryta.	Cauted strontia is insoluble. The Sinter, and assumes a black or corbonate, mixed with an equal volume of soda, fuses to a clear glas, which becomes milt white on cooling. In stronger hecomes cauting, and is absorbed by the coal.	Bintary and assumes a black of dark-gray color.
8· E	Constito time neither frace nor is it altered. The carbonate be- comes caustic, of white color, gives brightly, acquires alta- line reaction, and it a fragment be thus hestod, it falls to pow- der upon moistening with water. Readed that from the forces, the order fame acquires a failar-red coder.	Radiy soluble to a clear glass, that may be made opticated by faminar. The carbonate dissovers with effertresource. A larger addi- don gives a clear glass, which while cooling clear glass, which while cooling clear glass, which the and clouded, but never so milk-white as is the case with baryra and scronta.	Conside lime neither three nor is Radiy soluble to a clear giase, that Bohuble in large quantity (carbon. Insoluble; the code is absorbed by is perfectly infinible, and become altered. The carbonate be made operation by and the structure are which when considerably to each and isaves the lime on grows brightly, acquires alter the grass, which when considerably the surface. A larger addit giase, which when considerably the surface. A larger addit with the structure of structure of the structure of structure of structure of the structure of the structure of structure of the structure of the structure of structure of structure of the structure of structure of structure of structure of the structure of structure of structure of the	Tracitible ; the soda is absorbed by the cost, and leaves the lime on the surface.	la perfectly infusible, and become

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SHOWING THE BEHAVIOR OF THE

EARTHS, AND METALLIC OXIDES BEFORE THE BLOWPIPE.

Ķ.	The earbonate is decomposed : giows and soquires an alkaline resolica.	As itme, but is not so strongly arystalline.	be eachonais is decomposed i.A. Nme, but is not so strongly Discives easily (the outbornate.As Linne, giows and soquires an alkaline crystalline. resoltion. resoltion.		After long function semmers pais feath-red color, that must be ob- served when the assay is cold. Phosphate and arsenate of magnedia fuse and acquire a violet-red color.
8 Alterations	Unchanged.	Bowty soluble to a clear glass, which does not become opactuo, either by familing, or after com- plete saturation, by cooling, When added as a fine powder in large quantity, a glass results that is not clear, but on cooling becomes crystaline on the sur- tace, and is hardly fusible.	Blowly dissolves to a clear glass, that remains dear. With too large an addition the undissol- ed portion is rendered semi- trausparent.	Intumesces slightly, forms an intualitie compound, and the excess of soda is absorbed by the coal.	an After strong ignition becomes the beautifully blue, best observed when the assay is cold.
Gladina Ve	Unchanged.	Soluble in large quantity to a clear As with Boraz, base, that becomes milt-white by faming or when saturated, by simple cooling.	As with Boraz,	Insoluble.	Acquires a pale binish-green col- or.
Tutria. T and Tertia. Tr	Unchanged.	As Glucina.	As Glucina.	Insoluble.	0
. Brtula	The yellow ordde becomes of a lighter color in R. F., and ac- quires a transparent appear- ance.	ie becomes of a Dissolves slowly to a clear color. As with Borar, n R. F., and ac- less glass, which turns milk- usparent appear- white by flaming, or after satu- ration by mere cooling.	As with Borax,	Insoluble.	0
Zirconia Kr	Infusible. As propared from sul- As Glucina. plate, glows more intensely than any other substance.	As Glucins.	Dissolves more slowly than in Insoluble. Borax, and more readily yields an opaque glass.		Assumes a dirty violet color.
fi fi	Unshanged.	Boluble to small extent, forming a As with Borar. clear glass, which when fully clear glass, which when fully clear glass, which when fully controg; but if it remains clear after cooling, cannot be made opaque by flaming.	As with Borar.	Insoluble.	0
	Vincinged.	glowly soluble to a clear, difficult- ly fusible glass, that cannot be made opsque by fiaming.	Slowly soluble to a clear, difficult. To a very small degree soluble to With not too much soda, soluble With little colvait solution bocomes by funble glass, that cannot be a clear glass. The undissolved with effervescence to a clear faint in color. With more made opeque by flaming. The thinnest edges may, how eath.	With not too much mode, soluble with effervescence to a clear glass.	With little cobait solution becomes faint blutah in color. With more solution is ulack or dark gray. The thinnest edges may, how ever, be thesed by the stirtle blowing to a reddiab-blue stast.

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EARTHS AND METALLIC OXIDES.

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etallic Ox- es & Acids	Metaliko Oz. Ides & Acids Behavior alone, on Charcoal, etc.	With Borax on Platinum Wire.	With Salt of Phosphorus on Pla- tinum Wire.	With Bods on Cosl.	With Solution of Oobalt in 0. 7.
Antimon- eus acid 80.8	res, and mostly de d the basted point, et a again. A green- ge is communicated fiame.	O. F. Largely soluble to a clear plass, but becomes colories on cooling. On charcoal the dis- solved action any bealmost com- pletaly expelled from the beal, solved action as below. J. P. The glass which has been action as below. J. P. The glass which has been arrough the autor trunc to the O. F. on charcoal becomes gray- han and the glass become by contuned holowing these volatilize, and the glass become clear. With the the glass is used arrow to be a the glass is the arrow of the glass become sate is a less or more saturated.	O. F. Largely soluble to a clear of the solution to a clear plane, which while of a solution to a clear plane, which while of a solution of orders on other glass, which while connect other and sharity yellow. To there are also when ignated in O. F. is a solution of orders on other mediately volatilized, and not contrains while while and the other molecular plane, and solution of orders of the architectular plane, beam of the volatilisation of the architectular plane beam of the architectular plane beam of the architectular plane beam of the volatilisation of the architectular plane beam of the architectular plane architectular plane beam of the architectular plane architectular plane architectular plane architectular architectular plane architectular architectular architectular plane architectular plane beam of the architectular plane architectular architectu	In oharooal in both fames very easily reducible with the metal is firmediately volatilized, and covers the coal with a white de- posit of orde of antimony.	If the sublimate formed on ohar- coal when ignical in O. F. is bacted after moistaing with bacted after moistaing with solution of cobalt it is partly when the part of the second visities that and after cooling is seen to have acquired a dirty, dari- green color.
Amemous Acid	Volatilizes below a red heat.	0	0	On charcoal is reduced, evolving vapors of arsenic, which may be recognized by their garlic odor.	•
Oride of Burnuth N ³	O. F. On platinum foil fitaes easily to a dark-brown mass. which is pale yallow when cold. On charcoal in both flames it is re- duced to metalliob ismuth, which gradually voistilizes, depositing on the support an inter coating of yallow oxide, and nouter white one of carbonate of his- muth. In R. F. these sublimates disappear without thaging the flame.	0. F. Rasily solutile to a clear yel- low glass, which with a small quantity is coloriess when not, but while cooling becomes yellow, and is opaline when not, but while cooling becomes yellow, E. F. On charcoud the glass is at first gray and turbid, then the diret gray and turbid, then whe coids clear. The reduction co- cutes low. The reduction co- cutes low.	O. F. Eastly soluble to a clear yel. On tharcoal is immediately reductor when cold. With a karge quanta of F. on write dissolves to a tity of oxida, the glass can be bead, yellow when hot, pale yellow that the still more it becomes in 0. F. on write dissolves to a tity of oxida, the glass can be bead, yellow when hot, pale yellow that the still more it becomes low, opaque, when cold. With still more it becomes low, opaque, when cold. The f. F. for const espectation with help of the the glass so changes that concleas but on cooling becomes dark gray and opaque.	oluble to a clear rel. On charcoal is immediately reduce which is coloriess ed to metallic bismuth. With a large quan. In O. F. on wire dissofves to a With a large quan. In O. F. on wire dissofves to a with the gives can be beed, yellow when hot, pals yel- ue by faming, and more it becomes colling. It becomes colling the becomes bet it person clear a but on cooling be- gray and opeque.	•
Ortide of Oedminum Ód	O. F. On platfinum foil unohang- ed. R. F. On charcad disappears, and condenses on the surrounding cool as a redata, thrown to dark- yellow powder, the color of which is best seen when cold The exterior parts of the subil- mats are iridescut, like ithe fail of a pescock.	O. F. Boluble in very large quan- tity to a clear yillowiah glass, which is almost coloriess when oold. When strongly saturated the glass maybe made milt-wilt- by fiaming; very strongly satura- ied it becomes opeque by coloring. The cadmium is reduced, built the void the support with dark yel- low oxide.	0 4	F. Targely soluble, forming a 0, F. Insoluble, clear gass, which with a large 2, F. To charded a immediately quantity is yellowith which hock for charced. The metal volutions when nod, A sather and deposits reddish-brown and unsted glass is milk-white when the paid deposits reddish-brown and unsted glass is milk-white when nod. A sather and deposits reddish-brown and unsted glass is milk-white when nod. A sather and deposits reddish-brown and unsted glass is mulk-white when nod. A sather and deposits reddish-brown and unsted glass is mulk-white when nod. A sather and deposits reddish-brown and unsted glass is mulk-white when nod. A sather and deposits reddish-brown and unstered glass is mulk-white when nod. A sather and deposits reddish-brown and unstable glass is a sufficient of the sublimate is a sufficient of gluow oxide on the imate the seduction. The failure sufficient is reduction.	9

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TABLE OF BLOWPIPE REACTIONS.

Recording 1 a	rei gias (libs oxide of iron); on coulds, however, the bead is yellow. At a certain saturation the persones opeque to coulds. R. P. The pulow glass is randered colories. A highly artituted glass becomes enamel while and grave becomes enamel while and or F. Dissolves alowly, but with interse color. In small quarti- transformer and the pulow- when hot (chromic add) and when hot (chromic add) and when hot (chromic add) and when hot (chromic add) and when entirely cold fine yellow- hot, yellow while cooling, and when entirely cold fine yellow- is F. W. With ittle ohrome ordic the glass is besutful green hot, yellow when do and ond ; on a larger addition it becomes dark- to r pure emerald green the glass started and obt on contes. Starts which when euclidy saturated grow- entible trougs exturated grow- entible the network and when with the actively cold in yellow- the glass is a besutful green to chrome ordic on a contest gass, which when euclidy saturated grow- entible a nonorise. The network when the latter is added in larger the network and when when the latter is added in larger the network and assumes a hinthi- gray color. In selin green the seture of a columbility when the latter is addited in larger that the opedue glass form 0.7. Is uncharged, but with the owne of a seture of a building the antity the glass form 0.7.	A. Y. A. W. M. Downs: use over however, completely dilationable to an conting. Provide the providence of a contraction courses the galass to become opeque on cooling. A. F. Boluble to a clear glass to become opeque on cooling. A. F. Boluble to a clear glass to become opeque on cooling. A. F. Boluble to a clear glass to cooling. A. F. Boluble to a clear glass to cooling. A. F. Boluble to a clear glass to cooling. A. F. Boluble to a clear glass to cooling. A. F. Boluble to a clear glass to cooling. A. F. Boluble to a clear glass to cooling. A. F. Boluble to a clear glass to cooling. A. F. Soluble to a clear glass to cooling. A. P. A bound dirty green, and cool fing becomes dirty green, and clear glass the colors appear a some with thi. A. F. Soluble in large quantity the some solid. The lower oxide gives a fine glass is brown; addition of subast bulo, becoming brown is didition of subast bulo. D. F. With a large quantity the glass is brown; addition of subast bulo. Date of tron makes the lood red. The lower oxide gives a the cours divide the lower oxide gives a fine to blass becoming blass is brown; addition of subast bulo. Date of tron gives a blood-red oddr.	The product the sequencies of a light- gray color. Frank on the surface, of a light- gray color. 0. F. Soluble on plathum wire to a dark brownish-green glass which becomes opeque solor of and the control add). R. F. The glass is opeque and green (contob vero dot. On charcoal it remains as green oxide on the surface of the charcoal, while the soda it is absorbed. O. F. With about an equal yol- ume of the charcoal, while the soda it is absorbed. D. F. Asin O. F. The add can- be reduced to metal. D. F. Asin O. F. The add can- be reduced to metal.	While hot appears gray, on cool- ing become dirty gray, on cool- hag be too strong, the area. If the heat be too strong, the sean most strong heated are of a dark gray color.
O. F. Unchanged. R. F. Bhrinks together somewhat, and without fusing is reduced to metal, which is lifted by a magnet, and when rubbed in a mortar assumes metallic lustre.	O. F. The orde of cobalt possesses very great coloring power. The giass is mual tolline both when hot and cold. When strongly saturated the color is so deep as to appear black.	O. F. As with Borar: with equal quantity of orded the color is not quite so deep as with borar, es- pecially after cooling. R. F. As in O. F.	Q.F. On platinum wire dissolves in very alight quantity to a tran- parent, pads, roes of mass, that becomes gray on coording, that B. F. On charcoal reduced to a gray magnetic powder, which is made instrouts by rubbing.	•

METALLIC OXIDES.

Metallic Ox- ides & Acids	Metaliko Oz. Behavior alone, on Charcoel, etc.	With Borax on Platinum Wire.	With Salt of Phosphorus on Pla- tinum Wire.	With Bods on Coal.	With Bolution of Cobalt in O. P.
Ortic of Oopper On	 F. Fuses to a black globule, which on charcoal soon spreads out, and is reduced to metal on its under surface. R. F. is reduced at a temperature below the fusing point of me- tallio copper. The reduced por- tions have the metallic instree of copper, but as soon as the blow- ing is interrupted, the surface ordicals, By stronger heat the re- black. By stronger heat the re- black and bounds brown or black. 		0, F, A small quantity gives the 0. F. The colors are the same as 0. F. On platimum wire soluble to glass a green color with borax, but less intense; a clear green glass, which loses which charges quantity gives a green to diark or operus the color and bocomes opeque on fig. A larger quantity gives a green when hot, and blue or gooling. A larger quantity gives a green when hot, and blue or glass, which when hot, and blue or gooling. A larger quantity grees a green when hot, and blue or glass, which when hot, and blue or gooling. A larger quantity grees a green when hot, and blue or gooling. A larger quantity grees a green when hot, and blue or gooling. A larger quantity grees a green when hot, and blue or gooling. A larger quantity grees a green when hot, and blue or gooling. A larger guantity grees a green when hot, and blue or gooling. A larger quantity grees a green when hot, and blue or gooling. A larger the moment of a point grass and on cooling to the degree of saturation the R . F . A pretry strongly saturated grass soon becomes the opper and cooling the moment of a point grass becomes the opper and cooling at the moment of a point grass becomes colories in a grass of moment of a point grass per on glass becomes the transid with grass the moment of a cooper with duroed and separated so that the in on colling, a the moment of a cooper and opper with duroed and separated so that the duroed, it is onorise the source for the degree of saturated with grass of the duroed and separated so that the moment of a cooper and solvers. The duroed and separated so that the duroed at the source duroed at the grass of the source solvers at the moment of a cooper and solvers at the moment of a cooper and solvers with the moment of a source solvers at the moment of a cooper at a source of the source at a source at a source at a source of the source at a source of t	¹ F. The colors are the same as 0. F. On platinum wire soluble to with horax, but less internes; a clear green glass, which loes ris.; green to durk or opaque its colors and bue or coling. The colors and bue or coling. The order of a clear result reducting to the degree of saturated the to metallic copper, which there is a precise and super set or globules in a strong glass theoration. If the glass theoration and the theorated heat. A preting and the solution on coling, and the solution with the strong glass theoration. If the glass one is the nument of an in solution with but but becomes argue on while heat. If the glass one with the order of a copper the strong with no charced, it is colored as the solution with the order of copper the order of copper with the order of copper with a colores of a the definition of the structure of the order of copper the order of copper the order of copper with a structure of the order of copper the order of coppe	•
Oxfde of Didymium Di	O.F. Infusible, R.F. Losse its brown color and becomes gray.		O. F. Boluble to a rose red colored As with Borar, but more difficult. Insoluble. by the ci glass, which is unchanged in B. ly insoluble. F.	Insoluble. The soda is absorbed by the charcoal, leaving on its surface the oxide with a gray oolor.	o
Oride of Gold Âu ³	Ignited in either faume, is convert- ed into metal which is easily fusible to globules.	fame, is convert O. F. Is reduced without dissolv- As with Borax. I which is easily ing, and on charcoal may be ules. R. F. As with O. F.		As with Borar ; the soda, however, is absorbed by the charcoal.	o
Oxide of Ladium La	 F. Becomes dark yellow when heated, and igniter again on cooling. Intrusible. F. F. Gradnally reduced and valatilized, costing the coal and volatilized the outer flame violet. 	O. F. Becomes dark yellow when O. F. Dissolves to a clear glass, heated, and lighter again on feebly yellow while hot color- cooling. Intusible. Intusible. The set out of the set of th	As with Borar; but the glass when O. F. Insoluble. As with Borar; but the glass when O. F. Is reduced on coal, and the becomes gray and turbid on cool- ing the coal with orded, and ing.	O. F. Insolute. R. F. In reduced on coal, and the metal volatilizes in part, cost- ing the coal with orde, and party remains in the first in al- most aliver-white grains.	O
Oride of Iridium Ir	Is reduced by ignition, but the metallic particles cannot be tused.	gnition, but the O. F. Is reduced without dissolv- As with Borax, doles cannot be into piobules can be ob- tained. Z. J. As with O. F.	ds with Borac,	As with Borar ; the soda passes into the charcoal.	D
-7	للملاقية فسطان ملتدجني	an in the Billeria and indexes and and the fat in	a the first of the barrent of a		

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TABLE OF BLOWPIPE REACTIONS.

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magnetio powder.	Insoluble. The soda passes into the charcoal, and the oxide re- mains behind as a gray powder.	C. F. Eastly soluble to a clear O. F. As with Borar; more ortido O. F. On platinum wire easily yellow glass, which is a cooling; is rendered paque when old. Wile hot, are defined at a certain grade of while hot. The farming at a certain grade of while hot. The farming at a certain grade of while hot. The glass containing ordio for charcoal immediately returnated becomes operate and the glass containing ordio most and the glass containing ordio and becomes tracked and the glass containing ordio for the point of the maked to metallic lead, which seturated the far to be an ording. T. The glass containing ordio for the contact is metally becomes are and becomes fraction, and the glass point and the glass becomes turbid, but it is nover the glass becomes the far becomes the glass the gla	F. The glass requires much of F. On platinum wire slightly oride before it becomes concred, soluble to a clear green mass. When the bis becomes conclude with the management of the management of plate the management of oright, the management conclor, the management excellent. The special point of the solar is absorbed and the the mangement conclor, the solar is absorbed and building the hot bead in contact, with a green on color it, the the mangement excellent of the the mangement excellent of the the mangement excellent. The plate of the the mangement excellent of the the mangement excellent of the the mangement excellent. The object by as ordide.
If coloricat. With a very large quantity, the hot glass is dark red; on cooling it becomes brownish-red, then dirty green. The color disapter source by cooling than those of the borar glass. F. With little orded the color is not altered; with more it is red when hot, and on cooling first four, then greenish, and final- tin the glass becomes green and finally coloriess on cooling.	As with Borax	F. Eastly soluble to a clear 0. F. As with Borar; more orido or cooling, is rendered opaque is necessary, however, to pro- ory faming at a certain grade or while hot. By faming at a certain grade or while hot. The second solution of the second solution of the saturation, and still more highly R. F. The glass containing orde asturated becomes opaque and becomes grayth and turbid on errane! yellow on cooling. The glass containing orded orde, while harve does of apread islow on cooling. The glass containing orded order of the maken glass ued blowing the orde is reduced more turbid, but it is never with efferveecence to metallic entirely opaque.	0 4
Not, and yellow when Cold; with still more the hot glave is dark red, and when cold is dark yellow. <i>R. F.</i> The glass is bottle green (proto-sequicited). On charcoal with tin, is at first bottle green, then virtel green (protozide).	O. F. Soluble to a clear colorless glass, which at a clear colorless tion can be made opaque white by flaming, and more atrongly esturated becomes opaque by cooling. R. F. As in O. F.		O. F. Colors intensely. The hot glass is amethystine red, on cooling becomes violet red; with too large quantity the glass be- onnes opsque, and appears black, unless flattened, or drawn into threads. F. F. The colored glass becomes ovortess (protoxide). If the glass is deeply colored, the re- duction succeeds best on char- coal, especially with addition of tin.
	Unchanged.	Red lead heated on platimum-foil blackness and by gentle grantion is converted into yellow ordie: more strongy based this ordie frames to a yellow glass. On oharcoad in 0 , N , and R . P , is immediately reduced to metallo lead, which gradually volkitiles by continued heating, and ov- ers the oharcoal with a yellow deposit of ordie of lead, beyond which a thin which conding of actronate of lead is formed. These oscilage disposer when heated in R , P , tinging the fame same blue.	O. F. Infusible. The higher or- ides are converted by strong ig- nition into redich brown proto- sequioride, y- B. F. As in O. F.
angularida a Lroa Ha	Oxyd of Lantha- num	Ortice of Lond	Orida of Kangunan Hin

METALLIC OXIDES.

Metallio Ox- ides & Arida	Metaliko Oz- Behavior alone, on Charcoal, eto. idee & Arida	With Borax on Platinum Wire.	With Salt of Phosphorus on Pla- tinum Wire.	With Boda on Ooal.	With Solution of Oobalt in O. P.
Oride of Mercury. Hg	la immediately reduced and vola- tilized.	0	ò	Heated in closed tube (as well aloue) it is reduced, and con- denses in the coil parts of the tube as a graylah metallic sub- tube are which are to the tube the gray of the numbing with a feature, or better by cutting of the gray of the numbing the sublimate, placing it in a the sublimate, placing it in a the sublimate, placing it in a hydrochlorio sedi, and boiling tho latter.	o
Molybdio (Acid 1	O. F. Fusses with a brown color and volatilizes, condensing on the surrounding charcoal in form of a yellow sublinate, which near- est the assay consists of small crystals. The sublimate is white (the crystals colories) when cold. Interfor to this de- posit is een best when cold a thin, non-volatile, dark copper- ted coating of cold of a deep with the R. F. becomes of a deep with the R. R. P. becomes of a deep with the R. R. P. becomes of a deep with the R. R. P. becomes of a deep with the solution of the state of the the table model of the the table of the table be obtained by washing as a gray powder.	<i>F.</i> Fuses with a brown color and 0. <i>F.</i> Easily antible to a clear volating: conconsting on the surrounding charconsting on the surrounding charconsting on the surrounding charconstin form of low which with a moderate guartity of the assay is yould be assay in the hot, and atmost set the assay on the surrounding charconstin form of low which with a very large quantity of the assay is yould be assay is yould be assay in the non-transmitter of a solution of the assay on the distribution of low which mast. The multimate, which mast approximation of the assay on the distribution of the assay on the distribution of low which once the assay is proposed that which mast. With a very large onclustes of framily of the assay is produced the assay and through the low of the assay on the distribution of the assay is produced to the assay and through the low of the distribution of the assay to onclust of the assay is produced to the assay and through the low of the distribution of the assay and through the large on the distribution of the distribution of the assay the distribution of the assay and the distribution of the distredit distribution of the distrib	he brown color and (O. F. Zeady and largely soluble to a clear (J. F. On platimum wire fraces with contenting on the contention on the contention on the contention of the seasy is yellow with color but is colorated as the which hot brut is colorated as the moderate plane. The platimum wire fraces with color plane is a clear glass, which has the old. With a moderate plane, plane plane, which hot, but is colorated as the moderate plane is a clear glass, which has the old. With a work has the plane is a clear glass, which has the old. With a work has the plane is a clear glass, which has the old. With a work has the plane is a clear glass is dark yellower and almost cooling. The mail quarky the plass is dark yellower and almost cooling. The mail quark to plane the optime of a data almost cooling. The subminates is 10% to dark red when hor, and the glass becomes dark fractroscoled by the moltane optime or optime or optime of moltane is the subminates in the strongly staturated glass [R. F. The glass from O. F. be molybdio and mass is had and of the station dark to contact the support, and most of the station dark fraction of the station of the station dark to contact the support is plane to contact the station of the station of the station dark to contact the station of the state absorbed by the glass, when the later the state contact to contact the state. The state of the state is the state of the state	F. Hastly soluble to a clear O.F. On platimum wire fuses with grass, which with a molecute grass, which between the clear grass quantity of the assay is yellow which becomes in milk-which on the stress when odd. On thar A. A. On charvoal fusion with cooling. A. Y. On charvoal fusion with cooling. A. Y. On charvoal fusion with cooling. A. Y. On charvoal fusion with cooling. A stress of the support, and most of the support, and most of the molyhdio acid is reduced to cooling, however, fins green is a steled gray powder by walk-influence. With this, he green in coole somewhat darker.	0
Oride of Nickel Mi	R, F, On changed, R. P. On charcoal is reduced to metal. The coherent metallic product cannot be fused; strong. If reduces a metallic lustre, and is highly magnetic.		O. F. Colores intermedy. In small O. F. Dissolves to a reddiah glass, violet, which becomes the hot glass which becomes yealow on cool. R. F. On charcond setuly reduced to strong the becomes yealow on cool. R. F. On charcond setuly reduced the provided which becomes pale red in grass is brownish red, and the reduction provides reddiah yealow on cool. R. F. On charcoal setuly reduced argumutities these colores are dark. becomes reddiah yealow on cool. R. F. On charcoal reddiah yealow on cool. R. F. The glass is brownish red, and to reduce the comes gray and throw in the reddiah yealow on cool. R. F. The glass is provide the comes gray and throw it from the reddiah yealow on cool. B. F. The glass is three reddiah yealow on cool. R. F. The glass three contexts. Becomes reddiah yealow on cool. R. F. The glass three reddiah yealow on cool. R. F. The glass three reddiah yealow on cool. The glass is proved with the metallic for the reddiah proved with the sources are becomes at the reddiah proved are three. On separation of metallic nicel. Is explored the context on proceeds more reddiation of the glass becomes reduction proceeds more reddiation of the set with the nicked by the set of the set of the glass becomes reduction proceeds more reddiation of the set	 F. Insoluble. F. On charcool early reduced to granul brilliant metallito par- ticles, which are highly mag- netic. 	٩

TABLE OF BLOWPIPE REACTIONS.

0	o	0	σ	After long ignition appears light gray; on cooling, however, be comes plate red, line magnedia If it is not quite free from allali it sinters, and becomes bluich black.
Is early reduced to an infudble motalito powder, which may be obtained pure by washing.	Tasoluble. The soda is absorbed by the charcoal, leaving the Falledium behind as an intua- ble powder.	As Palledium.	is immediately reduced ; fuses to metallic globules, while the noda is absorbed by the charcoal.	In unchanged, except in color be 0. F. Easily soluble to a clear 0. F. Largely soluble to a clear 0. F. Mired with a little more After long ignition appears light coming family yellow when the more glass, which at a certain saturation of a solution when been accessed at a solution when more and a solution when more and a solution becomes on coloring. However, be the again on cooling. However, be the again on cooling family sature that an equal volume of a edga gray i on cooling. However, be the again on cooling and white again on cooling. However, be the again on cooling and white again on cooling and the again on cooling. However, be the again on cooling and the again on the cool at the cool and the again on the cool at the cool and the again on the cool at the again on the cool at the again of the again on the cool at the cool and the again of the
o	As with Boraz.	As Palladium.	and O. F. Both the oride and the Is immediately reduced; fuses to The metal yield a yellowiah glass. metallic globules, while the acda aline A highly saturated bead appears to the opsime on cooling it is color is yellow by transmitted idylight, be and red by candle-light, and arat-	O. F. Largely soluble to a clear glass, which if very highly satu- rated usy sollowiah when hot, and becomes coloriess on cooling. R. F. The glass from O. F. is un- changed.
c	In reduced on ignition, but the 0. F. Is reduced, without dissolr. As with Bornz. metallic particles cannot be fuse. Ing in the flux. The metallic particles cannot be united to a globule even on charcoal.	As Palledium,	metallic aliver, O. F. Is partly dissolved, and pobules. D. F. Is partly reduced to metal. The glass on cooling becomes opaline or milk-white, according to the degree of saturation. D. F. be- comes at first gray from sepa- ration of metal, then dear and cooless, all the silver separat- ing and fusing to a globule.	O. F. Eastly soluble to a clear glass, which at a certain satura- tion is pellowich when hot, be- comes colories on cooling, and may be made turbid by faming. At a greater degree of satura- tion becomes enamel white on cooling.
 P. F. Is converted into emails acid, which volatilise, yielding acid, which have a very pendering which have a very pendering and pargent odor, and attack if P. F. is reduced to a dark horwn furmal! to powder (metallic or infuelle) which may easily be oridized again to comic acid. 	Is reduced on ignition, but the metallic particles cannot be fus- ed together.	- As Palladitum.	Radiy reduced to metallic sliver, which tuses to globules.	la unchanged, ercopt in color be- coming faintly yellow when hot, and white again on cooling.
Outlds of Outlds of Os		Ortides of Platinum Fit Bibodium Hi Buthenium Hiu	Oride of Billrow	Tuntallo Acid. Ta

METALLIC OXIDES.

Metallio Or-	Metallio Or-Bohavior alone, on Charcoal, etc.	With Borax on Platinum Wire.	With Sait of Phosphorus on Pla- tinum Wire.	With Soda on Coal,	With Bolution of Oobalt in O. F.
Tellarous Acid Te	O. F. Fuses, and is reduced with effervescence. The reduced meta. volatilized however, im- mediately, and a white coating of tellurous acid deposits on the support. The deposits on the limate have commony a red or farme is tinged bluich green. flame is tinged bluich green.	F. Fuses, and is reduced with O. F. Soluble to a clear colories As with Borax. effertseence. The reduced with becomes gray from meta-volatilizes, however, in-geparation of metallic or the clinrium mediately, and a white conting when heated on charceal. F. The effect of no call becomes first filmate have commely a red or gray and finally colories all dark yellow color. The edges of the sub first elumin being cedical or colored for a state of the sub filmate have commely a red or the sub filmate have comments and costing filmate have comments are cost with tellmous acid.		On platfinum wire soluble to a clear coloriese glass, which be- comes withe on cooling. On charcoal it is reduced and vola- tilized with the formation of a costing of tailurous acid.	o
Binoride of Fin Bin		O. F. Very slowly soluble in small quarkity to a clear colorless glass, which remains clear after cooling, and is not made turbid by flaming. A bead saturated with oride, sulved to become perfectly cold, and then heaked nanifeste indistinct crystallisa- marifeste indistinct crystallisa- marifeste indistinct crystallisa- reaftered, a portion may be reduced.	 0. F. The protoride of the takes 0. F. Very slowly soluble in small 0. F. Very slow structure in solution in the interval slow of solution in the interval	O. F. On platinum wire unites with sola with effervesence, to a svollen infusible mass. R. F. On charosal is reduced to metallic tin.	Assumes a binish-green ooler, which must be observed after the assay is perfectly cold.
Ittanio Acid II	In both flames becomes yellow when heated; on cooling re- sumes its white color. Is not otherwise changed.	C. F. Easily soluble to a clear glass. The glass is yellow while not, coloress when cold, and not, bo rendered turbid by family, if it contains a large quantity of the assay. When the glass contains a very large the glass contains a very large the glass is yellow, the boomes opeque white on cooling. R. F. Dissolved in small quantity the glass is yellow to brown. A saturated glass may be made enamel biue by flaming.	becomes yslow (J. F. Bacily soluble to a clear (J. F. Radily soluble to a clear (J. F. On charcoal soluble with Assumes a yslowidh grees color, to colories when cold, and much of the subsection of the subsec	2. F. On charcool soluble with effervescome to a dark-relieve glass, which on cooling crystal- lizes with production of so much heat that that the globule becomes again of itself which bot. When fully cold the glass is while or. F. Ai in O. F. No reduction f. F. Ai in O. F. No reduction for metal can be accompliabed.	Assumes a yallowfah-green color, tamilar to orde of stor, but less fue.

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TABLE OF BLOWPIPE REACTIONS.

 J. Tochanged, unlies in very interne boat, when, as in R. F. it become black, being re- duced to tangedo oxida, but does not frue. 	O. F. Infutible, but is converted into durk velowinh-steen oxide. R. F. Becones black pasting into protoxida.	Furthle. The portions in contact with the charcond are reduced, and pass into the support; the remainder assume the color and instre of graphite, being con- verted into a lower oxide of vanadium.
O. F. Kaily sould so to a clear outoriess glass. Added in pretty hryse quantity it appears yolow while book with nore of the sub- stance the bead may be made ensure like by thaming, and with a still larger quantity it becomes opaque whito on cool- ing. F. The glass containing but little turgetio solf is unaitered in R. F., but as the quantity it of the solf is unaitered in R. F., but as the quantity is prove or darkyellow ool, and on cooling becomes yellowish- beson the subfactor. The same re- sections stocsed on charcosl with the art the color of tha glass when not too much tungsten is pres- ent.	O. F. Behavior like that of orde of from but the colors are less deep. When very strongly sat- urated, the glass may be made. In the grant of the grant of orde of from. The grant glass at a certain startation may be become enister enanel.itie nor crystalline. With the on char- coal the glass becomes dark- green (protorde).	ortions in contact (O. F. Soluble to a clear glass, O. F. Soluble to a cost are reduced, which is colorises with a small which when the quantity with more appears too small, has a muse the color and relaw, and on cooling becomes color while hot, and philo, while point are the color and the relaw and on cooling becomes pails yellow is hower code of E. F. The glass the chrome becomes fine chrome color and because the chrome color and the relax of the chrome code of E. F. The glass the chrome code chrome code of the chrome code of the chrome code chrome code chrome code chrome ch
F. Unchanged, unlies in very [O. F. Badiy soluble to a clear (J. F. Dapkinum when discreted is polyw which when as in hyse quantity it appears yellow highly attracted is yellow which when hyse quantity it appears yellow highly attracted is yellow which when hyse quantity it appears yellow highly attracted is yellow which when highly attracted is yellow and have here which when highly attracted is yellow while while while or yellow and may be reduced to matched to matched by the more other when odd it when highly attracted is yellow while here while hot. With an it alter quantity it have and in yellow at the more other when highly attracted is yellow at the more other highly attracted in the more odd the becomes opeque while on cooling becomes draft green when highly attracted is yellow or hor when highly attracted is yellow or hor highly attracted in the highly attracted is yellow or hor when highly attracted is yellow, highly attracted in the here of a data having a moduli of the farse of the highly attracted in the highly attracted in the here of a data having a moduli of the here of the matched in the here of a data having a moduli of the here of the matched with the except when here of a data having a moduli of the here of the matched when here of the matched when here of the matched here of the matched when here of the matched in the here of the matched here of the ma		The portions in contact O. F. Soluble to a clear giase, [O. F. Soluble to a clear giase, Puese with a suborted as the contact of F. Soluble to a clear giase, [O. F. Soluble to a clear giase, [O. F. Soluble to a clear giase, [O. F. Soluble to a clear giase, [D. F. Soluble to a lower code of R. F. The giase from O. F. As with Borax.
A.P. Chainum when classive to a lear dark-yellow glass, which an cooling becomes crystalline, and operate white or yellowith. R. F. With a lithe soda on char- coal may be reduced to metallio tungsten ; with more soda the sasy is absorbed into the ohar- coal, and yellow or brown tungs- tatio of soda having a metallio hustre is outsined.	.F. Soluble to a clear yellow O. F. Insoluble. With little soda glass, that becomes yellowiah. shows afgras of tusion; with grass from on coling. more soda tha mass becomes i. F. The glass from O. T. Bows afgras of the reagent, the assy pene- tore soluting. The grass from the reagent, the assy pene- ion of the grass from the reagent, the assy pene- however, is fine green (proto- trates the charcoal. Results of the reagent, the assy pene- bowever, is fine green (proto- darker (protoxide).	Fuses with soda, and is absorbed by the charcosi.
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METALLIO OXIDES.

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& Acids	Metallic Or-Behavior alone, on Charcoal, and des & Adds in the Platinum Tongs.	on Charcoal, and With Borax on Platinum Wire.	With Salt of Phoephorus on Pla- tinum Wire.	With Boda on Coal.	With Bolution of Oobalt in O. F.	-
Oride of Eino. En	O. F. Beormes yellow on heating, O. F. Easily and largely soluble As with Borax, but remnastiz white color when to a clear glass, which while hold on out. It is intrustible, and grows is yellowship, no considerably retricted and grows is yellowship, and when considerably reduces and distributions. When considerably represent the metal volatifier part deposite a control optically resturated may be made opeque and the highly estimated becomes and while when oold. R. P. The estimated becomes the highly estimated becomes and while when cold. R. P. The estimated becomes the start and while when cold. The host of the oxidely by longer blowing is considerably be oxide to pract the oxide hereal volatifies and while when cold. The oxide here al volatifies and deposites are over the exidely. The estimated when the start and while when cold. The oxide here al volatifies and deposites are over the exidely by longer blowing is considerably becomes the exident the exide here al volatifies and the export.	O. F. Eastly and largely soluble to a clear glass, which while hole is yellowish, on could becomes colorless. When could becomes asturated may be made opaque by familig, and when more highly esturated becomes highly esturated grass when first heated becomes turbid and first heated becomes turbid and the oxide). by longer blowing is the oxide), by longer blowing and the oxide), by longer blowing and dueed, the metal volatilizes and		O. F. Insolubie. R. F. On charcoal is reduced. Assume a fine yellowish-green The mediata, however, volatilises immediately, and if the best best estrong, burns with a bright greenish-white fame, while the charcoal is costed with orida.	Assumes a fine yellowish-green color, best observed when cold.	

TABLE OF BLOWPIPE REACTIONS.

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Chapter 3.

ALPHABETICAL LIST OF ELEMENTS AND COMPOUNDS,

WITH THE MOST CHARACTERISTIC BLOWPIPE AND OTHER REACTIONS EMPLOYED IN THE FOLLOWING TABLES FOR THE DETERMINATION OF MINERAL SPECIES.

96. Alumina. The only characteristic blowpipe reaction is the blue color it assumes when ignited with cobalt solution. It may be thus detected in most minerals of which it is a large ingredient, provided they are infusible and do not contain too large a quantity of colored metallic oxides, or of magnesia. Very hard minerals, like corundum, must be finely pulverized (79). From acid solutions, when neutralized with ammonia, alumina is thrown down as a flocculent white precipitate.

97. Ammonia. The slight green tinge that salts of ammonia impart to the blowpipe flame (76) is too faint and uncharacteristic to serve for their detection.

Ammonia is recognized by its well-known odor. The body to be tested is mixed with dry soda, the mixture placed in a closed glass tube, and gently heated, when the ammonia is evolved in the gaseous state, and may easily be recognized by its characteristic odor, as well as by the alkaline reaction it gives with readened litmus and with turmeric paper.

It must be borne in mind that organic substances containing nitrogen yield ammonia when ignited with soda.

98. Antimony. 1. Is almost invariably recognized by its characteristic sublimates. The body should be tested first in the open tube (74, 2, c); afterward, and generally in case of metallic compounds, on charcoal (75, 5, d).

2. Where antimony is combined with bismuth and lead, it is best detected by treating the substance with fused boric acid on charcoal, in such a manner that the flux is covered with the blue flame, and the metallic globule lies at its side partly out of the flame. The oxides of lead and bismuth are absorbed by the boric acid, and the charcoal becomes coated with a sublimate, which, when the blowing has not been too strong, consists of oxide of antimony, entirely free from the oxides of lead and bismuth.

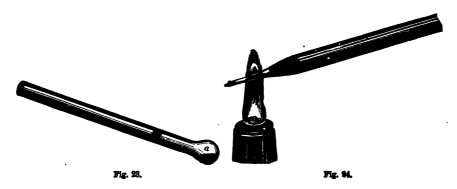
3. A small quantity of antimony, combined with copper or with other metals which retain it strongly, may volatilize so slowly that no sublimate forms on the charcoal. Under these circumstances, the alloy is heated in O. F. with a bead of salt of phosphorus, until the latter has dissolved a part of the antimony. The glass is then removed to a clean place on the charcoal and treated with tin in R. F. If the glass becomes turbid and black, antimony is indicated. Bismuth, however, gives the same reaction.

4. In examining sulphide of lead for antimony, compare 118, 2.

5. Compounds of antimony and arsenic, heated for a short time in the open tube, yield a mixture of crystals of arsenous acid and amorphous antimonous acid. A small amount of antimony mixed with sulphide of arsenic is detected by gently heating the dry mixture in a closed tube; the sulphide of arsenic volatilizes, while the dark-colored sulphide of antimony mostly remains where the assay was placed. The tube is then cut off between the two sulphides, and the sulphide of antimony is transferred to an open tube and tested as usual. When the quantity is extremely small the tube is crushed, and the fragments with adhering sulphide are introduced into the open tube.

99. Arsenic. 1. The testing in open tube (74, 2, a), closed tube (73, 11, f), and on charcoal (75, 5, c), usually lead to its detection.

Arsenous and arsenic acids and their salts, as well as the sulphides of arsenic, are examined by pulverizing and placing them in a glass bulb, covering them with six times their weight of a dry mixture of equal parts of cyanide of potassium and (arbonate of soda. The bulb should not be more than half filled with the mixture (Fig. 23). It is first gently heated; if moisture is given off, it is removed by inserting s. piece or roll of bibulous paper. It is again gently warmed, and if necessary viped out with paper, and the operation repeated until the mixture is *perfectly* dry. Finally, the bulb is heated strongly for some minutes in the spirit-lamp or lowpipe flame; a mirror of metallic arsenic deposits in the cool part of the tube. If the tube be cut off between the mirror and the sealed end by notching with a file and breaking, and the mirror be heated in the spirit-lamp, the arsenical odor will then be perceptible.



2. Arsenous acid can also be detected by introducing the assay into a closed glass tube drawn out to a small diameter (Fig. 24), and inserting a splinter of charcoal above it. The charcoal is first heated and then the assay; the arsenous acid is reduced as it passes over the hot charcoal and is deposited as in the previous case as a metallic mirror.

3. The higher arsenides, when treated in the open tube, yield a sublimate of arsenous acid, but the lower arsenides of nickel, cobalt, and iron do not part with their arsenic at a high temperature, even in the presence of reducing agents; and for its detection in these cases Plattner recommends the following method: Mix the finely divided assay with five times its weight of nitrate of potassa, and heat as intensely as possible in a platinum spoon. The metals are thus oxidized and the arsenic becomes arsenic acid. The spoon with the fusion is now boiled with water, until it is as far as possible dissolved. The liquid containing all the arsenic as arsenate of potash is decanted or filtered from the insoluble metallic oxides, and, 1. Evaporated with addition of a few drops of sulphuric acid (enough to expel all nitric acid) to dryness in a porcelain capsule; the residue is pulverized, mixed with cyanide of potassium and carbonate of soda, and heated as just described; or, 2. It is made slightly acid by acetic acid and boiled to expel any carbonic acid, and a crystal of pure nitrate of silver added, when a reddish-brown precipitate of arsenate of silver will be formed.

4. A small amount of arsenic in the presence of much sulphur is often difficult

to detect by its odor on charcoal. In such cases it is best to mix the assay with an excess of carbonate of soda, which will retain the sulphur, and the arsenical fumes can then be easily recognized.

100. Baryta. All the salts of baryta except silicates yield the characteristic yellowish-green coloration of the flame. When observed through copper-green glass the baryta flame appears bluish green.*

In Harmotome and Brewsterite, baryta is detected by dissolving the finely pulverized mineral in pure hydrochloric acid with aid of heat, filtering the solution and adding dilute sulphuric acid; a white precipitate of sulphate of baryta is formed, which may be collected upon a filter, washed, and then examined for the coloration of the flame.

The presence of the sul-Strontia may interfere with the baryta reaction. phate of baryta with the sulphate of strontia can be detected by fusing the mixture with three or four parts of chloride of calcium in a platinum spoon, and boiling the fused mass with water. If a cloudiness is produced, by adding to the clear dilute solution a few drops of chromate of potassa the presence of baryta is indicated. Strontia is only precipitated from the concentrated solution (Chapman).

101. Bismuth. 1. Bismuth is detected by the characteristic lemon or orange yellow sublimates which it and its compounds give when treated alone or with soda on charcoal in R. F. (75, 5, g). The presence of other easily oxidizable metals may make this reaction uncertain; the wet way must then be resorted to, and for this purpose the pulverized compound is digested for some time with hot nitric acid, the liquid poured off from any undissolved matters, or if necessary filtered, then evaporated almost to dryness, and the concentrated liquid poured into a test-tube half filled with water. If bismuth be present, a white precipitate of basic nitrate is formed, which may be collected on a filter, washed with pure water, and examined on charcoal. If the precipitate be small, it should be gathered into the apex of the filter; the latter is then dried, the part containing the precipitate torn off, and tested on charcoal.

2. If a compound of bismuth be treated with a mixture of equal parts of iodide of potassium and sulphur, and fused B. B. on charcoal, a beautiful red sublimate of the iodide of bismuth will be deposited.

3. In the presence of lead and antimony bismuth can be detected in the following manner : The mixture of the three oxides is added to an equal volume of sulphur and treated in a cavity upon charcoal with R. F.; the oxides are thus converted into sulphides. The assay is then placed upon a flat coal and treated with the O. F. and R. F. until the antimonial fumes have nearly ceased. The residue is placed in a mortar and pulverized, and mixed with an equal volume of a mixture of one part of iodide of potassium and five of sulphur; it is then heated in an open glass tube, and if bismuth be present, a distinct red sublimate of iodide of bismuth will be deposited a short distance above the yellow sublimate of lead. The sublimate of iodine which is liable to be deposited higher up the tube must not be confounded with the bismuth sublimate.

See also 98, 3.

102. 1. Borio (boracic) acid is recognized by the intense yellowish-green color it or its compounds with fluorine communicate to the flame. This color is given to the outer flame by most borates, provided they do not contain an ingredient which of itself tinges the flame.

* The strips of colored glass alluded to in this chapter are such as are used for colored glass windows, a cobalt-blue glass, a green glass colored either with oxide of copper or iron, and a red glass colored with red oxide of copper. Strips 3 × 6 inches are a convenient size. Von Kobell. Journal für Praktische Chemie '2), III. (1871), 469.

Corpwall. Am. Chemist, March, 1872.

2. Borate of soda alone tinges the flame pure yellow, but if it be moistened with sulphuric acid or mixed with bisulphate of potash, boric acid is set free, and the green color is instantly produced.

3. Silicates in which the above methods fail to indicate the boric acid, are reduced to a fine powder, the assay mixed with its own bulk of pulverized fluor-spar, and three times its bulk of bisulphate of potash; the whole is moistened to a paste, a portion of which is taken on a platinum loop, and at first gently heated to dry it, then more intensely in the edge of the blue flame. At the instant of fusion the green coloration appears, but is usually only momentary, so that the observer must direct his attention closely to the assay during the ignition.

4. As in the above trials copper and phosphoric acid may be mistaken for boric acid; it is sometimes best to use Rose's test with turmeric paper. To the solution of any borate hydrochloric acid is added until the liquid gives a distinct acid reaction (till blue litmus is reddened by it); a strip of turmeric paper is half immersed in the solution for some time, and the paper dried at a gentle heat (not over 212° F.). The smallest trace of boric acid gives the immersed portion of the paper a reddish-orange color. Silicates are fused with carbonate of soda in a platinum spoon, the mass is boiled with water until it is as far as possible dissolved, the solution is then supersaturated with hydrochloric acid, and tested as above.

The orange or reddish-orange color thus produced must not be confounded with that communicated to turmeric paper : 1st. By alkaline solutions. 2d. By acid solutions of zirconia (159). 3d. By moderately strong hydrochloric acid.

5. If alcohol is poured over a borate with the addition of a sufficient quantity of concentrated sulphuric acid to liberate the boric acid, and the alcohol kindled, the flame, particularly on the edges, appears of a very distinct yellowish-green color, especially upon stirring, and upon heating the alcoholic mixture.

103. Bromine. 1. When bromides are added to a bead of salt of phosphorus which has previously been saturated with oxide of copper, and the blowing continued, the bead becomes surrounded with a beautiful blue flame inclining to green on the edges, and this color continues so long as any bromine remains. As these reactions may be confounded with those given by chlorine, Berzelius recommends fusing the substance under examination with dry bisulphate of potash in a glass bulb. If a metallic bromide is present, bromine and sulphurous acid are set free, and the glass bulb becomes filled with a yellow vapor of bromine, which, although mixed with sulphurous acid, may be distinctly recognized by its characteristic odor. As a confirmatory test, if moistened starch or starch paper be exposed to these vapors yellow bromide of starch will be formed.

2. If a soluble bromide be placed upon a piece of clean silver along with a fragment of sulphate of copper or sulphate of iron, the silver becomes almost immediately coated with a black stain.

104. Cadmium. This metal can only be detected as oxide, as it is volatilized at a comparatively low temperature. The substance for examination in spulverized state is heated in the R. F. on charcoal, whereby metallic cadmium is volatilized, and immediately on coming in contact with the atmosphere is converted into oxide which gives the characteristic coating on coal (75, 5, j). Should the substance contain not more than one per cent. of cadmium, as for instance in many zinc ores, it is best to mix the powder with soda and heat carefully in the R. F., when the coal near the assay becomes coated with a sublimate of **oxide** of cadmium before any sublimate of zinc is formed, cadmium being much more volatile than zinc.

Caesia. This rare alkali imparts a beautiful violet to the blowpipe flame, and when mixed with potassa and rubidia can only be distinguished by the employment of the spectroscope. 105. Carbon and Carbonic Acid. 1. *Carbon* in the form of diamond or of graphite, disappears when heated for some time B. B.; the former leaves no residue, the latter generally more or less of a red ash.

Fused with nitrate of potassa, carbon detonates, forming carbonate of potassa. *Carbonates* effervesce when treated with dilute hydrochloric acid; a few require to be pulverized, and in some cases heat is necessary before the effervescence * takes place.

2. Some carbonates lose their carbonic acid by simply heating in the closed tube; in these cases it may be detected by inserting a strip of moistened litmus paper in the tube, when the blue color will be changed to red, but on drying the original blue color will be restored.

3. Organic substances, except oxalates and formates, decompose in the closed tube, yielding a burnt odor, and usually oily products. *Anthracite* gives off moisture, but no empyreumatic oil. (See *Coal*, in the tables, chapter iv.)

106. Cerium. When in combination with other earths, cerium cannot with certainty be detected B.B. In most silicates where it, with lanthanum and didymium, occurs in considerable quantity, it may be readily detected after separation of silica and precipitation by ammonia, by treating the washed ammonia precipitate with oxalic acid, which dissolves out iron with alumina, leaving the cerium earths as insoluble oxalates; this residue when washed and ignited gives a cinnamon-brown powder, which is the characteristic color of sesquioxyd of cerium.

107. Chlorine. 1. Chlorides, like bromides, may be detected by adding a small portion of them to a bead of salt of phosphorus which has previously been saturated with oxide of copper; the bead becomes instantly surrounded with a beautiful and intense purplish-blue flame, without any of the tinge of the green which is observed in examining a bromide.

2. The soluble chlorides give the same reaction as described under bromine with sulphate of iron and copper on a silver plate.

3. Nitrate of silver produces, even in highly dilute solutions of hydrochloric acid or metallic chlorides, white curdy precipitates of chloride of silver, which upon exposure to the light change first to violet and then to black.

108. Chromium. 1. Chromium is detected by the emerald-green color which its compounds impart to the borax and salt of phosphorus beads. Chromium must not be confounded with vanadium, which gives the same reactions in R. F., but differs by yielding a yellow bead with salt of phosphorus in O. F., which flux never acquires other than a green color from chromium.

2. Minerals containing but little oxide of chromium associated with other metals which color the fluxes, are best treated by fusing on platinum wire or in a platinum spoon with a mixture of equal parts of soda and nitre. The mass is heated for some time in O. F., whereby chromic acid is formed. The fusion is dissolved in water, and the solution poured off from the residue; to this solution a drop or two of acetic acid, and afterward a crystal of acetate of lead, are added, when a lemonyellow precipitate of chromate of lead is formed. This may be collected on a filter, washed, and tested with borax and salt of phosphorus.

3. A mineral which contains a small amount of chromium, and is not decomposed by nitre, is fused with one and a half times its volume of soda and three-fourths its volume of borax to a clear bead; this is pulverized, dissolved in hydrochloric acid, and evaporated to dryness, dissolved in water; the residue of silica filtered off; the protochloride of iron changed to sesquichloride by boiling with a few drops of

[•] Care must be taken not to confound minerals which contain a carbonate as an impurity with pure carbonates. If the substance under examination be a pure carbonate it can be completely dissolved in nitric acid, and effervescence will continue so long as any portion remains undissolved.

nitric acid, and the chromium, alumina, iron, etc., precipitated with annonia. The precipitate is collected, and tested as above.

109. Cobalt. 1. In most cases can be recognized by the characteristic blue bead it gives in both flames with borax. This color is variously modified by other metals.

2. Should iron be present, the glass will appear green while hot, and blue when cold. If the substance contains copper or nickel, the cobalt-blue color can hardly be perceived, and the bead must be treated on charcoal, with tin in R. F., until it becomes transparent, and effervescence has ceased. The copper and nickel will be reduced to the metallic state, and the glass will have a perfectly pure blue color.

3. Compounds of cobalt with arsenic, and arsenides of other metals, when fused upon charcoal until arsenic fumes cease to be given off, then treated with borax in R. F., give, when freed from iron, a pure smalt-blue color; if iron be present it will be oxidized before the cobalt, and the bead will have a bottle-green color. The metallic globule is then treated with a fresh quantity of borax, and this operation is repeated until the bead gives a pure cobalt reaction.

. In testing metallic nickel for cobalt it is necessary to combine the nickel with arsenic, which may be done by mixing the finely divided nickel with metallic arsenic, placing it in a depression in the charcoal, and fusing in R. F. The fused globule is then tested with borax, as just described in case of an arsenide. The volatile metals in combination are recognized by their sublimates on charcoal.

110. Columbium. If a mineral which contains columbic acid be powdered and fused with bisulphate of potassa, the fused mass powdered and dissolved in water, the columbic acid, and tantalic acid if present, will be insoluble; while the bases and titanic acid, if present, will be dissolved, and can be thus separated. The residue is treated with sulphide of ammonium, to free it from tungstic acid and oxide of tin, if these be present, and after filtration and thorough washing it is treated with dilute hydrochloric acid to remove traces of iron. The residue is treated with hydrochloric and sulphuric acids, with the addition of metallic zinc. If only a tantalate be present, no coloration ensues, or but a slight one. If a columbate is similarly treated, the separated columbic acid rapidly assumes a blue color, which gradually fades, and finally becomes brown.

111. Copper. 1. The green color which most copper compounds give to the blowpipe flame, and the reactions of its oxides with the fluxes, render its presence easily detected. The production of a red bead with salt of phosphorus in R. F. is rendered more certain by the treatment of the bead on charcoal with a small amount of tin.

2. Copper may also be detected by saturating a salt of phosphorus bead with the substance containing it, and adding chloride of sodium, when the bead will color the flame beautifully blue, owing to the formation of chloride of copper.

Many minerals give this reaction by simply moistening in hydrochloric acid and exposing in the platinum forceps to the flame; silicates should be first pulverized, moistened with hydrochloric acid, and evaporated to dryness in a porcelain capsule; then made into a paste with water, and heated on platinum wire.

3. In case the copper is combined with nickel, cobalt, iron, and arsenic, the greater part of the cobalt and iron may be separated by treating with borax on charcoal. The remaining metallic globule is fused with a small quantity of pure lead, and then boric acid is added; this last dissolves the lead and the rest of the cobalt and iron, while most of the arsenic is volatilized. The cupriferous nickel globule, which still may contain a little arsenic, is treated with salt of phosphorus in O. F.; the bead obtained will be dark green while hot and clear green when cold. This last green is caused by a mixture of the yellow of oxide of nickel and the blue of oxide of copper.

4. According to Guericke,* a very delicate test for copper is to mix the substance under examination intimately with chloride of silver, and fuse on iron wire; in this manner the smallest quantity of copper may be detected by the blue color imparted to the flame.

112. Didymium. See p. 36.

113. Erbium. See Yttria.

114. Fluorine. 1. Hydrofluoric acid imparts to Brazil-wood paper a strawyellow color. Silicates containing even a small quantity of fluorine, when heated in the closed tube, give off hydrofluo-silicic acid; this is decomposed into silicic acid, which is deposited near the assay and hydrofluoric acid, which passes off, and the latter may be detected by inserting a strip of moistened Brazil-wood paper at the open end of the tube.

2. When fluorides are heated in a glass tube with bisulphate of potash, hydrofluoric acid is given off. This etches the tube immediately above the assay, and gives the reactions with Brazil-wood paper just mentioned.

3. The best method for the detection of fluorine in all cases is to mix the assay with previously fused salt of phosphorus, and heat in the open tube in such a manner that the flame passes into the end of the tube.

In this way hydrofluoric acid is formed; it may be recognized by its peculiar pungent odor and its corrosive action on the inner surface of the glass tube, rendering it opaque and lustreless at the points where moisture has condensed. For a confirmatory test the reaction with Brazil-wood paper may be employed.

As the heat required in this experiment is so great that the glass tube often becomes soft and unmanageable, it has been recommended to use a piece of platinum foil rolled together and inserted into the end of the glass tube, as in Fig. 25.



FIG. 25.

The substance to be tested is placed with the flux upon the projecting part of the foil, and the flame directed as before.

115. Glucina gives no reactions which admit of being determined B. B. with certainty (see page 33). It is not of frequent occurrence, being only found in combination with silica and alumina.

116. Gold may usually be recognized by its physical characters. It is separated from the easily volatile metals by simple heating on charcoal in O. F. If associated with copper or silver, it must be fused with a large excess of metallic lead and subjected to cupellation (see 142). The copper becomes absorbed and passes off with the lead, while the silver remains alloyed with the gold. If the globule is quite yellow it is proof that but little silver is present; it is then to be tested with salt of phosphorus to prove the presence of silver, which after fusion will impart an opaline appearance to the cool bead. If it be more of a silver color, the amount of gold will be small, and in order to prove its presence the globule must be digested with hot nitric acid in a test-tube or porcelain capsule; the **silver** is thus dissolved, and the gold remains in a fine powder or as a spongy mass. If this powder be washed and fused with borax on charcoal it will yield a globule of metallic gold. In combination with infusible metals, such as platinum, iridium, palladium, and rhodium, the alloy obtained B. B. is less fusible. For their separation the wet reagents must be employed.

117. Indium. Colors the flame beautiful violet. (See table, p. 36.)

* Pharm. Centralblatt, 1855; 195.

118. Iodine. 1. Iodides, added to a bead of salt of phosphorus which has previously been saturated with oxide of copper, tinge the outer flame an intense emerald-green color. (Compare bromine and chlorine, 103, 107).

2. Iodides, like bromides, are decomposed by fusion with bisulphate of potash; free iodine is liberated, and may be distinguished by its characteristic violet color and its disagreeable odor. If an iodide be added to a mixture of carbonate of lime and caustic lime, then intimately mixed with a small quantity of chloride of mercury and heated in a closed tube, iodide of mercury will be sublimed; this is easily recognized by its first yellow and then red-yellow color. It is best to draw the tube out to a narrow neck a short distance from the assay, and for the success of the experiment it is necessary that all the substances employed be perfectly free from moisture. This test is said to be even more delicate than the starch test, which is used in the wet way.

119. 1. Iron is distinguished by the characteristic color its oxides impart to borax and salt of phosphorus, as well as by its compounds yielding a magnetic powder with soda on charcoal. (See treatment with soda, 94.)

2. In the presence of easily reducible metals, such as lead, tin, bismuth, antimony, or zinc, iron may be detected by treating the assay with borax and charcoal in R. F., until everything except the iron has been reduced, when the borax glass will have a bottle-green color. If the substance contains much tin, or if the bottle-green glass is fused with tin-foil in R. F., the iron becomes entirely reduced to protoxide, and the bead has a pure vitriol-green color.

3. In case the substance contains cobalt, nickel, and copper, the two latter will be reduced by the tin, while the cobalt will color the bead blue. To detect the iron it is only necessary to heat a portion of the blue bead, with addition of fresh borax, on platinum wire in O. F.; the bead will be green while hot and blue on cooling.

4. To distinguish the presence of protoxide of iron in minerals, Chapman recommends the following method:—"A small quantity of black oxide of copper is dissolved in a bead of borax and platinum wire so as to form a glass which exhibits, on cooling, a decided blue color. To this the test-substance in the form of powder is added and the whole is exposed for a few seconds, or until the test-matter begins to dissolve, to the point of the blue flame. If the substance contain protoxide of iron it will be converted into sesquioxide at the expense of some of the oxygen of the copper compound, and opaque red streaks and spots of red oxide of copper will appear in the glass, as the latter cools. If only sesquioxide of iron is present, the glass on cooling will remain transparent, and will exhibit a bluish-green color.

120, Iridium. (See p. 36.)

121. Lanthanum. (See page 37.)

122. Lead. 1. Compounds of lead give globules of metallic lead when heated with soda on charcoal B. B. It is recognized by its physical properties, as well as the characteristic coating it gives upon the coal (75, 5, h). The coating is modified by the presence of various other volatile metals. In the presence of zinc, the characteristic color of the lead coating is recognized on cooling, since the oxide of zinc becomes white. In the presence of bismuth, the oxide of which often obscures the lead, it is detected by heating the sublimate in the R. F., when the flame will be tinged with the azure-blue color which is characteristic of lead in the absence of selenium. The presence of selenium in such cases is evident from its odor.

2. Combinations of sulphide of lead with other metallic sulphides are tested for lead by treating in the R. F. either alone, or with borax to separate iron; and the lead is recognized by its coating. In such combinations the oxide of lead is surrounded by a white coat of sulphate of lead, which renders the presence of small amounts of antimony uncertain. The safest way under such circumstances is to mix the powder of the substance with soda, which reteins the sulphur; this mixture, when treated in the R. F., gives the pure lead coat, and if antimony is present it is detected by its white sublimate beyond the sublimate of oxide of lead.

3. In solutions of the salts of lead, sulphuric acid gives a white precipitate of the sulphate of lead, which is nearly insoluble in water and dilute acids. It is best to add a considerable excess of dilute sulphuric acid, evaporate the solution on a water-bath, and add water to the residue, when delicate tests are to be made in the wet way.

123. Lime. Lime imparts a characteristic yellowish-red color to the flame. When observed through copper-green glass the lime flame appears siskin-green; with cobalt-blue glass it is pale greenish-gray, and is almost entirely obscured. Many lime salts react alkaline to test papers after ignition. It is distinguished from baryta and strontia in the wet way, by the fact that sulphuric acid gives no precipitate in dilute hydrochloric solutions. Sulphuric acid gives a precipitate in the concentrated solution which distinguishes it from magnesia.

124. Lithia. The red color which pure lithia salts give to the flame is more or less modified or entirely obscured when mixed with other substances. Seen through green glass the lithia flame appears orange colored, with red glass is colored deep red, but with cobalt glass of sufficient thickness the flame is invisible. Silicates containing only a little lithia scarcely color the flame red; but if the pulverized mineral be mixed with one part of fluor-spar and one and one-half of bisulphate of potash, the whole made into a paste with a little water and exposed on platinum wire to the point of the blue flame, the outer flame will be colored distinctly red. Chapman has proved that the lithia flame, unlike strontia, is not obscured by the presence of baryta. He suggests fusing lithia minerals with chloride of barium; the phosphate, triphylite, when thus treated gives a beautiful crimson color.

125. Magnesia is recognized by its reaction with nitrate of cobalt (80, 2).

In combination with other earths, the wet way must be employed for its detection. Sulphuric acid does not produce a precipitate in its concentrated solution.

126. Manganese. The reactions of manganese with the fluxes are so peculiar and delicate that it may be recognized even when it exists in the smallest quantity, and in the presence of almost every other substance. The soda test in O. F. on platinum wire is the most delicate. If a reaction be not obtained with soda alone, a small fragment of nitre should be added to the assay, and the mass again heated. When testing substances which do not dissolve readily in soda it is well to add a little borax to the bead, and this also makes the test much more delicate (Chapman).

127. Mercury and amalgams give a sublimate of metallic mercury when heated in a closed tube. Compounds of mercury heated in a closed tube with soda yield metallic mercury, which condenses on the tube above the assay. When a gray sublimate is obtained, without distinct metallic globules, the part of the tube coated with it is cut off and boiled in a test tube with a little dilute hydrochloric acid; by this treatment the mercury collects into shining globules. In case mercury exists in so small a quantity that the sublimed metal is not perceptible, it may be detected by inserting a piece of gold-leaf held on the end of an iron wire into the tube, just above the assay; on heating, the mercury is volatilized and unites with the gold, giving it a white color.

128. Molybdenum. The sublimate which molybdic acid gives on charcoal (75, 5, m) and its reactions with borax and salt of phosphorus serve to distinguish it in most instances.

When it is present in small quantity, particularly when associated with copper and tin, as in some furnace products, it is necessary to have recourse to the wet way. The solution of a mineral containing molybdenum in hydrochloric acid, or the hydrochloric solution of the fusion with nitre and soda of an insoluble substance, when boiled with tinfoil is colored dark blue by the separated molybdate of molybdenum (compare Tungstic Acid, 153). Molybdic acid can also be recognized by heating the finely pulverized substance in a porcelain dish with concentrated sulphuric acid, and then adding alcohol. The fluid when cold acquires a fine azure-blue color, especially upon the sides of the dish.

129. Nickel may be recognized by the color its oxide imparts to borax and salt of phosphorus, together with its easy reduction to the metallic state in R. F.

Arsenical compounds of nickel, cobalt, iron, and copper are treated with glass of borax (see Cobalt, 109). When the borax is no longer colored blue from cobalt, but acquires a brown color, which is violet when hot, the metallic globule is separated from the borax, and treated with salt of phosphorus in O. F. If copper as well as nickel be present in the assay, the glass thus obtained will be green both while hot and cold; treated with tin on charcoal it will become red and opaque on cooling. A small quantity of nickel occurring in cobalt compounds cannot always be detected by the foregoing method. In such cases Plattner recommends saturating one, or if necessary several borax beads with the substance on platinum wire. The beads are then fused on charcoal in R. F. with from 50 to 80 milligrammes (0.75 to 1 grain) of fine gold; the oxide of nickel, together with a small portion of the cobalt, is reduced to the metallic state and unites with the gold. The metallic globule is then freed from the flux and treated on charcoal in O. F. with salt of phosphorus. The bead itself will be colored blue, as cobalt is easier oxidized than nickel, or perhaps if a little nickel be also oxidized it will be dark violet while hot and dirty green on cooling; in both cases the globule is separated from the flux and treated with a new portion of salt of phosphorus. If the original bead with borax was not too saturated, this second bead with salt of phosphorus will be of a pure nickel color. Should copper as well as nickel be present in the gold globule, the salt of phosphorus bead will be green while hot, and retain its green color on cooling; treated with tin in R. F. as before described, the bead will become red.

130. Nitrates. When nitrates are fused in a glass tube with bisulphate of potash, dark reddish-yellow nitrous fumes are evolved. The color is best observed by looking lengthwise through the tube held against a white ground.

All nitrates detonate when heated on charcoal; those of the alkalies and alkaline earths detonate violently, and are converted into carbonates.

131. Osmium. See p. 39.

132. Oxygen. Oxygen is evolved from some compounds by simple ignition. The substance under examination is placed in a closed tube with a bit of charcoal above it, the charcoal is first brought to ignition, and then the substance is heated, when, on liberation of oxygen, the ignited splinter of coal will glow with increased brilliancy.

133. Palladium. See p. 39.

134. Phosphates. 1. The green color (76, 4, f) which phosphates give to the flame serves in many cases for their detection. This coloration is heightened by the addition of a drop of concentrated sulphuric acid, but is rendered unsatisfactory in the presence of other substances giving a green flame.

2. If a pulverized phosphate is fused in a closed tube with a bit of metallic magnesium or sodium, the phosphoric acid will be reduced, and if the fused mass on cooling is moistened with water, phosphoretted hydrogen will be given off, tecognizable by its characteristic disagreeable odor.

3. When a few drops of neutral or acid solution containing phosphoric acid are poured into a test tube filled to the depth of an inch with a solution of molybdate of ammonia with nitric acid, there is formed in the cold or after a short time a pulverulent yellow precipitate of phospho-molybdate of ammonia. The reaction is hastened by very gently warming, care being taken not to heat above blood heat. A yellow coloration of the fluid must not be regarded as proof of the presence of phosphoric acid, since silicic acid produces a strong coloration, but it does not give a precipitate. Arsenic acid gives the same reaction.

135. Platinum. See p. 39.

136. Potassa may often be detected by the violet color it communicates to the flame. In presence of other bodies that tinge the flame, especially soda and lithia, this reaction is masked. The potash flame when observed through cobalt-blue glass * appears purple, and may thus be easily detected even in the presence of lithia and soda. With green glass it is colored azure blue, and with 'red glass deep red.

In presence of soda, potassa may be recognized by fusing borax with addition of a small quantity of boric acid on platinum wire, then adding enough oxide of nickel \dagger to make the glass brown when cold; the substance is dissolved in the bead thus obtained; if potassa be present it will be of a beautiful blue color on cooling. With soda alone a brown bead will be obtained.

For the detection of potash in compound substances it is often necessary to have recourse to the wet way. Bichloride of platinum produces in the neutral and acid solutions of the salts of potassa a yellow crystalline heavy precipitate of the platinchloride of potassium. Very dilute solutions are not precipitated by this reagent, hence they should be evaporated before testing; or better, evaporate to dryness after addition of the reagent and then dissolve the residue in alcohol, in which the platinchloride is insoluble.

137. Rhodium. See p. 39.

138. Rubidia. This rare alkali gives B. B. a violet flame, and when mixed with cæsia and potassa can only be distinguished by spectroscopic examination.

139. Ruthenium. See p. 39.

140. Selenium. The reaction for selenium on charcoal (75, 5, a) is so characteristic that the slightest traces of it can thus be detected.

Selenites and selenates are reduced to selenides on charcoal in R. F. with the characteristic odor of selenium.

141. Silica. 1. When silica is heated with soda, a clear glass is obtained if the soda be not in excess. This reaction distinguishes silica from the earths; silica may, however, contain alumina and still fuse with soda to a clear glass.

In most silicates the silica may be detected by help of salt of phosphorus (see p. 26). The experiment should be performed with a small fragment, from which the bases will be dissolved, while the skeleton of silica will maintain the same form as the original assay and float about in the bead. Only when a fragment is unaffected the powder is used, but when thus tested the result is less satisfactory.

2. When a finely powdered silicate is fused with an excess of carbonate of soda, the resulting mass dissolved in dilute hydrochloric acid, and evaporated to dryness, the silica is rendered insoluble; and on moistening the residue with strong hydrochloric acid, and dissolving in hot water, the silica will remain behind, and can be separated from the bases if desired.

3. Most of the hydrous silicates, and many which are anhydrous, but which con-

• The blue glass should be of sufficient thickness to entirely obscure a lithia flame; there is no objection to using two or three thicknesses of glass if necessary.

+ Oxalate or carbonate of nickel (emerald nickel) may be employed. It must be free from sobalt (not give a blue glass with borax).

tain an excess of base, are decomposed by strong hydrochloric acid; the bases then unite with the hydrochloric acid, while the silica separates either as a gelatinous hydrate, or as a non-gelatinous powder.

142. Silver. Silver is recognized by its physical characters as well as by the brown coating it gives when heated on charcoal in O. F.

When associated with volatile and easily oxidable metals, it may be separated by heating on charcoal in O. F. If the silver be associated with a large quantity of lead or bismuth, it is best to subject it to cupellation. The following process serves for the detection of silver in most argentiferous minerals: The substance is mixed with its own bulk of borax glass and an excess of pure lead (except in cases where lead or its oxide already exists, as in litharge, minium, cerusite, etc.), the mixture is placed in a cylindrical cavity in the charcoal, and fused in R. F. The flame should at first be directed entirely upon the borax glass; after the earthy substances have been dissolved and the metallic particles united into one globule. this globule is subjected for a short time to the O. F., thereby separating such volatile and easily oxidizable substances as may be present. The remaining globule containing a large excess of lead and all the silver, together with the larger portion of the nickel and copper, is then separated from the flux and subjected to cupellation.

For this purpose finely pulverized bone-ash is mixed with a small quantity of soda, and made into a stiff paste with water. This paste is placed in a circular cavity in charcoal, half an inch in diameter and one quarter inch deep, and the surface of it made concave and smooth by pressing it with an agate pestle or other suitable convex surface. This *cupel* is now carefully exposed to a gentle heat till perfectly dry.

The lead globule, freed from all adhering flux, is placed upon the cupel, and treated in O. F. Should much nickel or copper be present, an infusible coating is formed which prevents the desired oxidation; this may be counteracted by the further addition of a small quantity of pure lead. The blast is kept up until all traces of lead have become oxidized; this is indicated by the cessation of the rainbow-colors of the oxide of lead which play over the surface of the button. When the quantity of litharge that is formed in the process of cupellation is large, the globule of silver, still containing lead, may be removed to a fresh cupel and there refined. The instant when the last traces of lead disappear can then be more readily perceived; this point is indicated by the sudden brightening of the globule. The remaining metal, when free from gold, has a silver-white color. It may be tested for gold as described under that metal.

143. Soda. Soda is readily distinguished even in compound substances by the intense yellow color it imparts to the outer blowpipe flame. The soda flame is invisible when observed through cobalt-blue glass and red glass; with green glass it is orange colored. Soda is not precipitated from solution by bichloride of platinum.

144. Strontia. The crimson color imparted to the outer flame serves in most instances for the detection of strontia and its salts. In the presence of lime this reaction is less characteristic, and a small amount of soda obscures it altogether. The color is intensified by moistening with hydrochloric acid. When the strontia flame is observed through cobalt glass it appears of a pale purple to rose-red color, through green glass it is orange, and with red glass it has a deep red color.

After ignition its salts give an alkaline reaction on test paper, and it is distinguished from lime, which also gives a red flame, in that its dilute solutions are precipitated after some time by sulphuric acid.

145. Sulphur. Sulphuric Acid. Free sulphur fuses and sublimes; on charcoal burns with a blue flame, forming sulphurous acid. The higher sulphides give off sulphur when heated in a closed tube; the neutral sulphides and sub-sui-

phides give off sulphurous acid when heated in an open tube. 'The sulphurous acid may be detected by its odor or by its reddening and bleaching action on a strip of moistened blue litmus paper. Small quantities of sulphides and the sulphur in sulphates may be detected by fusing with two or three parts of soda on charcoal in R.F. In using this test it should be kept in mind that illuminating gas often contains sulphur; where this is the case a candle or lamp flame should be used (see 93). The sulphur is hereby converted into sulphide of sodium, which, placed on a clean silver surface and moistened with water, causes a brownish or black stain on the silver. In the presence of selenium this reaction cannot be used.

The soda used for the detection of sulphur should always be tested by itself for sulphur, which is a common impurity, and if it give the reaction, it should be treated as described on page 30.

The solution of a sulphate in hydrochloric acid gives a precipitate of the sulphate of baryta, on addition of chloride of barium.

The following is a delicate test for sulphides in the wet way. An amount of the assay powder that can be taken upon the point of the knife is mixed with a like volume of iron powder (*ferrum alcoholisatum* of the apothecary), the mixture placed in a cylinder of glass two and a half inches long and about an inch in diameter, and hydrochloric acid is poured upon it (one volume concentrated acid and one volume water). A strip of filter paper, which has been moistened with acetate of lead and again dried, is placed beneath the cork that fits the tube, which is then closed, the paper projecting from the tube a short distance. In about one minute the color of the paper is observed, and the glass shaken if necesmary. If sulphur be present the paper will be blackened by the formation of the sulphide of lead.

146. Tantalum. See Columbium, 110. 147. Tellurium. 1. Tellurides heated in the open glass tube, give a white or grayish sublimate, fusible B. B. into colorless or nearly colorless drops. On charcoal they give a white coating, and color the R. F. green.

2. When a substance containing tellumium is triturated with soda and charcoal dust and fused in a closed tube, then allowed to cool, and a little hot water dropped into the tube, the water assumes a beautiful purple color from the dissolved telluride of sodium.

3. Tellurium compounds when gently heated in a matrass with much concentrated sulphuric acid, impart to it a purple color, which disappears on the addition of water, while a blackish-gray precipitate is formed.

148. Terbia. See Yttria.

149. Thallium. Colors the flame intensely green. (See p. 18.)

150. Thoria, gives no reactions which permit its determination with certainty.

151. Tin. In the metallic state, tin is easily distinguished by its physical characters and its reactions in O. and R. F. on charcoal (75, 5, l). Sulphides containing tin must be roasted, and the roasted mass treated with a mixture of soda and borax in R. F.; the product is metallic tin, which can be further tested on charcoal. Oxides containing tin are best treated with soda or cyanide of potassium on charcoal; if much iron is present borax should be added. When tin and some of its compounds are treated with nitric acid, oxide of tin separates as a white precipitate, which can be separated and tested as above.

152. Titanium. 1. The violet color given by titanic acid with salt of phospho rus in R. F. serves in most cases for its detection. In the presence of iron the violet color first appears when the bead is treated with tin in R. F. on charcoal.

2 If a substance containing titanium is fused with carbonate of soda, and the

resulting mass dissolved in hydrochloric acid, and then heated with tin or zinc, the titanic acid is reduced to sesquioxide of titanium, coloring the liquid violet, and finally the violet hydrated sesquioxide separates.

When the fusion of a substance with six or eight parts of bisulphate of potassa is dissolved in a very little water, the clear solution decanted from the insoluble residue and a few drops of nitric acid and five or six volumes of water added, titanic acid if present will separate on boiling as a white precipitate.

153. Tungsten. Tungstic acid gives a blue color with salt of phosphorus in R. F.; with much iron the bead becomes dark red, but treated on charcoal in R. F., with tin it gives a blue color.

When a tungstate is fused with carbonate of soda and treated with hydrochloric acid and zinc as above (see Titanic Acid), a fine blue color is obtained.

Tungstic acid is insoluble in acids; hence if a tungstate like scheelite is decomposed by acids, the tungstic acid separates as a yellow powder.

154. Uranium. The reaction with phosphorus salt serves in most instances for its detection.

155. Vanadium. In the absence of other colored metallic oxides, vanadium may be detected by borax and salt of phosphorus; it may be distinguished from chromium by the color which it gives to salt of phosphorus in the O. F.

156. Water. Water may be detected by heating the assay in a matrass or closed tube, care being taken to free the tube from all moisture before inserting the assay. If a substance contains hygroscopic water, or if it be a soluble hydrous salt, the water is almost immediately given off and condenses in the upper part of the tube in distinct drops. Insoluble substances containing water require to be heated somewhat higher. See further under examination in the closed tube, 73.

157. Yttria. (Erbia and Terbia.) For the detection of these rare earths recourse must be had to analysis in the wet way.

158. Zinc. The reactions of this metal on charcoal, together with the green color which the oxide gives with cobalt solution, allow of its being detected when it exists in considerable quantity-and even in extremely small quantities, if it be not associated with other metals whose reactions are such as mask those given by the zinc. If a small quantity of zinc be associated with large quantities of lead, bismuth, or antimony, it is with difficulty detected. If a mixture of different metallic oxides be fused with a mixture of two parts soda and one to one and a half parts borax, zinc will be volatilized, and in the moment of coming in contact with the air, is oxidized and gives a coating on the coal. If the substance contain a large amount of lead, this is also oxidized and coats the coal, but on moistening with cobalt solution and heating in O. F., the lead coating is reduced by the charcoal, and the zinc coating becomes green on cooling. If the quantity of zinc is extremely small, it is best to moisten the coal with cobalt solution before heating the assav. In the presence of tin and antimony it is almost impossible to detect small quantities of zinc B. B.

159. Zirconia. This earth as usually obtained gives out an exceedingly brilliant light when heated B. B. A dilute hydrochloric acid solution of zirconia, or of minerals containing zirconia, imparts an orange-yellow color to turmeric paper when it is moistened with the solution.

Chapter 4.

TABLES FOR THE DETERMINATION OF MINERAL SPE-CIES BY MEANS OF SIMPLE CHEMICAL EXPERI-MENTS IN THE WET AND DRY WAY.

TRANSLATED FROM THE TENTH EDITION OF FRANZ VON KOBELL'S "TAFELN ZUR BESTIMMUNG DER MINERALIEN."*

Introduction to the Tables.

THE object of the following Tables is to facilitate the determination of mineral species. By means of a few simple experiments before the blowpipe and in the wet way, the mineral is quickly limited to a group of a few species; among the members of this group the mineral is distinguished by other trials, and when from these various experiments the mineral species is finally decided upon, the conclusion is confirmed or corrected by reference to the physical characteristics given in the columns upon the right, and further confirmatory evidence may, if necessary, be obtained by reference to a treatise on mineralogy. An acquaintance with the use of the blowpipe, such as is gained by the study of the preceding pages, and with the manner of performing the simplest operations of solution and precipitation, is all that is necessary in making the requisite trials.

It is hoped that this little work will be of service to chemists, miners, and others, who though not making mineralogy a special study, yet have occasion to decide upon the names of minerals.

The Tables are so constructed that it is necessary to follow them through from the beginning, comparing the characteristics of each group and division with those of the specimen in hand. A trial of fusibility, a fusion with soda, heating the pulverized substance with acid, and a few precipitations, usually lead to the desired object, when the order of the Tables is strictly followed, and the experiments are made with proper care.

The method which has been adopted in the arrangement of these Tables will be comprehended at a glance. The minerals are arranged in two great groups, *metallic* and *non-metallic*, under which heads are various classes, divisions, subdivisions and sections, the more general ones being placed upon the left, until finally we reach more specific characters, followed by the names of the species, in the middle of the page, while the remaining columns are devoted to the confirmatory evidence of color, streak, cleavage, fracture, hardness, specific gravity, fusibility, and crystalline form. In a few cases these physical characteristics are the distinguishing features of the species, but generally the mineral will be recognized by its blowpipe and chemical reactions, and the student is strongly advised to make these primary, since the chemical composition of the minerals is what is desired to

• This chapter includes, essentially, all the material contained in the tenth edition of Professor Von Kobell's Tables, but an entirely different mode of arrangement is here given, with much additional matter. The tabular form in which the minerals are arranged was suggested by Professor W. T. Roepper, of Bethlehem, Pa., who kindly permitted me to consult a manuscript translation made by him from one of the earlier editions of Von Kobell, in which a similar arrangement is employed. The Tables here presented, while following the general idea of Professor Roepper as to tabulation, have been worked up independently, and contain new features which it is hoped will be of service to the student. be known, and naming minerals from their color or other physical properties often leads to serious errors, especially with inexperienced observers, and these alone need the caution, since the experienced person well knows the impossibility of always recognizing minerals from the evidence of sight. It is thought by this arrangement of the Tables that more definite ideas of the groupings of minerals will be gained by the student, and that he will more readily comprehend which are the general and which the specific reactions of the smaller divisions. Almost all the established mineral species are included, but for the sake of convenience, their relative importance, or frequency of occurrence, or facility of determination, has been indicated arbitrarily by the size of the type in which the name of the species is printed

An attempt has been made, as far as possible, so to arrange the groups and divisions, that such errors of observation as are likely to be made, shall not prevent one from arriving at a correct conclusion. Since some minerals occur in one variety with metallic, and in others with non-metallic lustre, and since the fusibility of a mineral often varies, or may be underrated or overrated by the experimenter, and since the constituent elements of some mineral species are not constant, such are found under **both** or all of the divisions to which they might be assigned.

The following general directions may serve to assist in the use of the Tables:-*Lustre.* Under the head of metallic lustre only those minerals are included which are perfectly opaque. To determine this a fine splinter or thin edge should be held between the eye and the light, or fine fragments should be placed upon a white plate, when, if the slightest translucency is observed, it is included under "non-metallic." It is evident that opacity alone does not make metallic lustre, but that the mineral must also possess the lustre which suggests it to be metallic, and must not grind to an earthy powder as do some non-metallic minerals which otherwise might be called metallic. In this, as in many other determinations, good judgment in the operator will be constantly required.

Fusibility. For determining the fusibility of minerals, the following scale is employed :

Scale of Fusibility.

1. Stibnite (antimony glance)	(large magments.
2. Natrolite	Fusible in the flame of a candle, in
3. Almandine Garnet (alumina-iron- garnet	Infusible in the candle flame, but easily fusible B.B., even in somewhat large
4. Actinolite	Fusible B.B., in rather fine splinters.
5. Orthoclase	Fusible B.B., in finer splinters.
6. Bronzite	B.B. becomes rounded only on the finest points and thinnest edges.

Splinters of these minerals are kept ready for use, and in determinations their fusibility is compared with that of like splinters of the assay. The evidence of fusion is the rounding of sharp edges. It should be remembered that some minerals swell up before the blowpipe but do not fuse, and other phenomena take place which without careful observation might be mistaken for fusion. Only the O. F. should be used, since some substances, which are infusible in the O. F. are easily fusible in the R. F., on account of the reduction of some of their oxides to a lower fusible state.

Hardness.—In testing hardness, the scale proposed by Mohs, and almost universally adopted, is here employed.

Scale of Hardness.

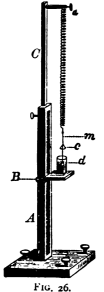
1. Talc.	3. Calcite.	5. Apatite.	7. Quartz.	9. Corundum.
2. Gypsum.	4. Fluorite.	6. Feldspar.	8. Topaz.	10. Diamond.

The scale represents the crystallized varieties of the minerals mentioned. The hardness of a mineral is found by finding what numbers will scratch, and what are scratched by the mineral to be tested. Thus, if a mineral will not scratch apatite, but will scratch fluorite, it is of a hardness between 4 and 5; or if the mineral is scratched by apatite and not by fluorite, it is of a like hardness. Sharp corners must be used in scratching, and particular care should be taken in this as in all other cases, that impurities do not come in to modify the result; thus a grain of sand in some of the impure varieties of galena, if it happen to come upon the corner which is used, would make the mineral appear quite hard, and without proper caution many such errors will be made.

Color.—Great care must be taken in forming any conclusions from the color of minerals. In minerals of metallic lustre, the color is generally constant, and often very characteristic, in some of the non-metallic species the same is true; but experience will teach how greatly the colors of non-metallic minerals vary, and varieties are constantly found differing in color from all that were previously known. Hence, especially in non-metallic minerals, the color which is given should only be regarded as an aid or suggestion in the determination.

Streak.—The streak of a mineral is tested by scratching it with a knife or file, or better, if not too hard, it may be drawn across a piece of unglazed porcelain, and the color of the mark which it leaves behind observed.

Specific Gravity.—Considerable skill can be gained by noticing the comparative weight of minerals held in the hand, and though no accurate determination can thus be made, the column giving specific gravity can be used in the field, as designating whether minerals are heavy or light. For accurate determinations, the apparatus, described in the foot-note below,* gives very quick results and in



* The specific gravity of minerals is easily taken by means of an instrument devised by Prof. Jolly. See Fig. 26. This consists of a graduated strip of looking-glass set in a vertical rod (A) properly supported. A steel or brass wire in the form of a spiral is suspended from a, and bears upon its lower end the two pans, c and d. The spring can be placed at any desired height by elevating the smaller rod (C), as shown in the figure. The pan d is suspended in water in the glass, which rests upon the sliding support B. At m is a signal which serves as a mark for the stretching of the spiral. The reading is taken by bringing the mark and the image of the mark in the mirror scale to a level. The scale d being in the water, the position of the mark m is taken = x. A fragment of the support B moved downward till the instrument again comes to rest, the scale d being still in the water, and the position of the mark m again taken = y. Then y-x = weight in air. The fragment is now transferred to d and the position of m again noted = z. Then y-x = loss of weight in write the scale is graduated is of little importance if it be regular, and hence the apparatus is easily constructed. This spring balance is known in Germany as the *Federwage*, and is furnished by Mechaniker Berbaride

most cases with advantage, can be made to replace the or linary chemical bal ance.

Testing for Water.—In order to detect water, a fragment of the assay is placed in the bottom of the closed glass-tube or matrass and heated strongly. Water, if present, condenses in drops on the cold part of the tube. A trace of moisture will be found by heating almost any mineral in this way; a little practice enables one tc decide whether or not the mineral is actually hydrous. Decrepitating minerals may be enveloped in a piece of copper foil, and thus placed in the tube and heated.

Decomposition by Acids.—In testing whether a mineral be decomposable by hydrochloric acid, it must first be pulverized as finely as possible in a mortar, and then gently boiled with tolerably concentrated acid for ten minutes or more, unless the solution is sooner completed. The digestion is carried on in a small glass flask, a large test tube, or a casserole. In cases where the fact of the decomposition is not evident to the eye, by the formation of a jelly, disappearance of the powder or other effects, the acid must be separated by decantation or filtration from the residue, ammonia or carbonate of soda added in excess, and then a few drops of phosphate of soda. When both these reagents give no precipitate, or cause but a few flocks to appear, the mineral may be pronounced nearly or quite undecomposable. The production of a decided precipitate is evidence that it has been decomposed.

Gelatinization. When silicates are decomposed by hydrochloric acid, the silicates sometimes separates in the pulverulent condition, when the mineral is said to be soluble in acid with the separation of silicates without forming a jelly; sometimes the silicates separates from the bases in its soluble condition, and then when the solution is boiled nearly to dryness, it will have the consistency and appearance of jelly. Such minerals are said to gelatinize with hydrochloric acid.

Many silicates not appreciably attacked by acids gelatinize after they have been previously ignited, as for example garnet, vesuvian, etc. Several splinters or little pieces of the assay are fused or strongly ignited, then pulverized and boiled in a test tube with rather dilute acid; on evaporating the solution gelatinous lumps will be seen in the remainder; or after standing some time (twelve hours) an evident fixed jelly will be found. After adding water, and stirring with a glass rod, the solution may be tested for bases if desired. Other silicates, which gelatinize with acids or are easily decomposed, will not gelatinize or are but little affected by hydrochloric acid after ignition.

Pyro-electricity. Some minerals when heated become electric, and have the power of attracting light substances. Light fibres of wool or cotton, or a deer's hair held between the fingers, may be used to test this property.

The methods for all the other commonly recurring reactions will be found under their proper heads in the preceding chapters.

In seeking the name of a mineral it is necessary always to begin with the first group and proceed in regular order to those following; for it often happens that a mineral belonging to one group has also the characters of the succeeding ones, while the minerals of the latter divisions may not show the reactions of the earlier groups. The same rule is of the greatest importance in the distinctions between subdivisions and species. Upon page 63 is given a summary of the classification—this is merely introduced to save turning the pages, and to give a more definite view of the larger subdivisions.

The method of using the table is best learned by some examples, first, without the use of the general classification.

Aluminite.

It is not metallic, turning over the pages which are headed minerals with

metallic lustre, we come on page 72, to the minerals without metallic lustre, to which group our mineral belongs. Looking now in the column on the left we see A.-B.B. easily volatile or combustible, which our mineral is not; looking along this column there follows B.-B.B. fusible from 1-5, etc. Our mineral is infusible; turning over to page 89 we come to C-infusible or fusible above 5. Looking in the next column we see that the members of DIVISION 1 are characterized by giving a blue color when moistened with cobalt solution and ignited; the mineral upon trial is found to belong here. This group is seen to be divided into two sections; in the minerals of the one water is present, in the other it is absent. By heating the specimen in a closed glass tube it yields much water; it must, therefore, be sought in section a. The minerals of the first sub-section give, on fusion with soda, a sulphuret which blackens silver, and since the assay gives this reaction it belongs here. Of the minerals which belong to this sub-section, the first is insoluble in hydrochloric acid and the others are not; on trial the powder of the mineral easily dissolves and it is, therefore, aluminite. Looking now in the columns on the right the determination may be substantiated by a comparison with the physical properties there tabulated. The chemical constituents and generally the formulas of the minerals are given, and should always be noted, so as to fix in the mind the composition of the various species. The formula for aluminite is $Al_2SO_6 + 9Aq$. In our examination we have detected all three of its ingredients : the alumina by the blue color with nitrate of cobalt; the sulphuric acid by the fusion with soda, and the water in the closed tube; but when in the determination of a species all the constituents are not determined, those who possess the requisite knowledge can, if desirable, detect the remaining substances by the ordinary methods of mineral analysis.

An example showing the use of the general classification will now be given.

Bornite (variegated copper).

Looking at the classification on page 63 the metallic lustre of the mineral places it under I. It is not a malleable metal. It is fusible and therefore belongs under A. B.B. it gives no odor of arsenic or selenium; gives no white coating which colors the R.F. or other reaction for tellurium, gives no fumes of antimony, but gives the reactions for sulphur, mentioned in division 5, and hence belongs to this division. We are now referred to page 67 on which this division is seen divided off in the second column. Looking now in the third column, it is not malleable, it gives no reaction for manganese, its streak is not red, it gives no globule of lead with charcoal (turn to next page), but moistened with hydrochloric acid it gives to the flame the blue color of chloride of copper, and it forms a sky-blue or green solution with nitric acid, which becomes deep violet-blue on addition of an excess of ammonia; of the minerals which give this color the first give a bismuth reaction, on trial the assay does not, but it fuses to a brittle steel gray magnetic globule, it hence belongs among these minerals. It has not the brass-yellow color of the first three minerals but has the variegated shades of the next, and is therefore bornite.

The ordinary varieties of mineral coal are included in the tables (see page 96). It hardly need be again remarked, that only pure and homogeneous material will give satisfactory reactions for the determination of minerals. If it is believed that the material being tested is not pure, regard must be paid to the impurity, and the reaction judged of accordingly; as, for example, many specimens of *wollastonite* (tabular spar) effervesce in acids, and after ignition impart a brownish red color to moistened turmeric paper. These qualities do not belong to the pure mineral, but come from an admixture of calcite. Too great haste should not be exercised in deciding upon the name of a mineral, since oftentimes the difficulties in the way of an accurate determination can only be overcome by long and careful labor.

On beginning the study of determinative mineralogy, it is best to examine known species, until confidence is gained in one's ability and accuracy. The following minerals are given by Von Kobell to his students; when these have all been determined, the student will be prepared to determine any mineral which can be distinguished by this method :--

> Aluminite, Alunite. Anhydrite, Antimony-Glance, Apophyllite, Argentite, Arsenopyrite, Atacamite, Barite, Borax, Bornite, Bournonite, Calamine, Calcite, Cassiterite, Celestite, Cerussite, Chalcopyrite, Chalcocite. Cinnabar, Cobaltite, Cryolite, Cuprite, Datolite, Diallogite, Dolomite, Fluorite, Galenite, Glaucodote, Gypsum, Hematite,

Lapis-lazuli, Lievrite, Lepidolite. Limonite, Magnesite, Magnetite, Malachite, Manganite, Molybdenite, Natrolite, Niccolite, Orpiment, Pectolite. Psilomelane, Pyrite, Pyrolusite, Pyromorphite, Pyrrhotite, Realgar, Scheelite. Smaltite, Smithsonite. Siderite. Sphalerite, Strontianite, Talc, Witherite, Wolfram, Wollastonite. Wulfenite.

(Page 63)

ANALYTICAL TABLE

SHOWING

GENERAL CLASSIFICATION

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MINERALS.

ABBREVIATIONS USED IN THE TEXT OF THE TABLES.

Amorph .. Amorphous. B. B.....Before the Blowpipe. DivDivision. Fib.....Fibrons F. Fus...Fasibility. Gran....Granular. H.....Hardness. HCl.....Hydrochlorio acid. Infus....Infusible. KI.....Iodide of potastium. Mass.....Massive. B......Near.

O. F..... Oridising fiame, p. c. Per cent. R. F.... Reducing fiame. S.... Sulphur. Sp. Gr... Specific gravity. Stalacc... Stalactito. I...... Isometric. II. Tetragonal. III. Hexagonal. IV...... Monocilinic. VI...... Trictinic.

A barred letter signifies two of an element; dots over a letter signify oxygen, as \overline{A} means Al_2O_3 .

GENERAL CL

PAC

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1.-MINERALS WITH METALLIC LUSTRE.

(Of these minerals whose lustre may be doubtful, only such are here included as are perfectly opaque on the thinnest edges. The native malleable metals and mercury are easily distinguished from others (see p. 64). The remaining minerals form the following groups.

A.—Fusible from 1-5, or easily volatile.

1.	B. B. on charcoal give the strong garlic odor of arsenic	(
2.	B. B. on charcoal or heated in an open glass-tube give the strong horse-radish odor of selenium.	ŧ
8.	B. B. on charcoal give a white coating which colors the R. F. green and greenish-blue. In a small test-tube, <i>gently</i> heated with much concentrated sulphuric acid, impart to the acid a hyacinth-red color, which upon addition of water disappears, and a black gray precipitate of tellurium is thrown down	
4.	B. B. on charcoal, or in the open glass-tube, give dense antimony fumes	E
5.	Heated in the open glass-tube, give sulphurous acid, which reddens a strip of moistened blue litmus paper placed in the end; or, with soda, give a sulphur reaction, but do not give the reactions of the preceding divisions	
6.	Not belonging to the foregoing divisions	E
	B.—Infusible or fusible above 5, and non-volatile.	
1	P. P. in O. F. give to the how a head the emotheration and of menomenon	r

•••	D. D. m. O. F. give to the bolax beat the antenystine fet of manganese	4
2.	Are magnetic or B. B. on charcoal after long heating in R. F. become so	7
8.	Not included in the foregoing divisions	1

II.-MINERALS WITHOUT METALLIC LUSTRE.

AB. B. easily volatile or combustible.	
--	--

B.—B. B. fusible from 1-5, and non-volatile, or only partially volatile.

Part L.-B. B. with soda on charcoal give a metallic globule, or fused alone in R. F. become magnetic.

	magnetic	
1.	B. B. with soda on charcoal give a globule of silver	7
2.	B. B. with soda on charcoal give a globule of lead	7
8.	Moistened with hydrochloric acid give a beautiful blue color to the blowpipe flame, and give with nitric acid a solution which, on addition of an excess of ammonia, becomes violet-blue (copper)	
	a) B. B. on charcoal evolve a strong arsenical odor	7
	b) B. B. on charcoal evolve no arsenical odor	7
4.	B. B. impart a beautiful sapphire-blue color to a borax bead (cobalt)	7
5.	B. B. fused in the forceps or on charcoal in R. F. give a black or gray magnetic mass, but do not give the reactions of the preceding divisions	
	a) During fusion evolve a strong arsenical odor	71
	 b) Soluble in hydrochloric acid without leaving a perceptible residue, and without gelatinizing 	7(

SIFICATION.

	AGE 77
	78
Not belonging to the foregoing divisions	79
Part II. —B. B. with soda on charcoal give NO metallic globule, or fused alone in R. F. do not become magnetic.	
B. B. after fusion and continued heating on charcoal or in the forceps have an alkaline reac- tion, and change the color of moistened turmeric paper to red-brown*	
a) Easily and completely soluble in water	80
b) Insoluble or difficultly soluble in water	81
Soluble in hydrochloric acid, some also in water, without a perceptible residue; the solution is not gelatinized by evaporation	82
Soluble in hydrochloric acid, forming a stiff jelly upon evaporation	
a) B. B. in the closed tube give water	83
b) B. B. in the closed tube give no water or but traces	84
Soluble in hydrochloric acid, leaving a residue of silica without forming a perfect jelly	
a) B. B. in the closed tube give water	80
b) B. B. in the closed tube give no water or but traces	81
Slightly attacked by hydrochloric acid, and B. B. give a deep amethystine color (manganese) to the borax bead	87
Not belonging to the foregoing divisions	87

C.-Infusible or fusible above 5.

First ignited B. B., then moistened with cobalt solution, and again ignited assume a beautiful blue color (alumina)	
	00
a) B. B. in the closed tube give water	09
b) B. B. in the closed tube give no water or but traces	90
Moistened with cobalt solution and ignited B. B. assume a green color (zinc)	91
After ignition B. B. have an alkaline reaction and change the color of moistened turmeric paper to red-brown	91
Vearly or perfectly soluble in hydrochloric or nitric acid without gelatinizing or leaving a con- siderable residue of silica	92
lelatinize with hydrochloric acid, or are decomposed with the separation of silica	
a) B. B. in the closed tube give water	93
b) B. B. in the closed tube give no water or but traces	94
Not belonging to the foregoing divisions	
a) Hardness under 7	94
b) Hardness 7 or above 7	96

* Kenngott has shown that many allicates and other compounds before and after fusion have an alkaline reaction when are placed upon turmeric paper in the form of possion and moistened with water; but they do not show this reaction when agments.

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(Page 64)

MINERALS WITH METALLIO LUSTRE.

NATIVE METALS.

A. Fusible from 1-5, or easily volatile.

Devasion 1 (in part),

1. MINERALS WITH

		General characters.	Bpecific characters.	Species.
		Alone, B. B., on charcoal, a bismuth coating.	With sulphur and iodide of potassium gives a red coating (bismuth), and leaves a glo- bule of gold.	
	લ્વઇર્સેઝે.	Soluble in nitric acid; the di- lute solution gives a precipi- tate with hydrochloric acid.	The precipitate becomes violet-gray on exposure to light.	Silver.
4	nau.	Have more or less the color of gold.	Only soluble in aqua-regia without a residue.	Gold.
Native Metals	their		Decomposed by aqua-regia with separation of AgCl.	Electrum.
	ed by	Of a copper-red color.	Moistened with hydrochloric acid colors O. F. sky-blue.	Copper.
Nati	Eastly recognized by their malleability	Give a lead coating on coal.	Easily fusible; soluble in nitric acid; the solution gives a heavy precipitate with sulphuric acid.	Lead.
	Eastly :	Infusible. Insoluble in hydro- chloric acid ; soluble in aqua- regia.	Insoluble in nitric acid.	PLATINUM
	•		Soluble in nitric acid.	Palladium.
		Attracted by the magnet.	Infusible. Soluble in hydrochloric acid.	IRON.
		Compare the malleable minerals, <i>Hessite</i> , Div. 3, p. 66; and <i>Argentite</i> , Div. 5, p. 67.		
		Fluid.		MERCURY.
		B. B. volatile without fusion.	Gives in the closed tube a metallic sublimate.	ARSENIC.
		With soda on coal give a glo-	Very fusible; gradually volatilizes; gives no copper reaction.	Dufrenoysi
	હ	bule of lead. Soluble in ni-	Decrepitates strongly.	Sartorite.
DIVISION 1.	charcoal give the garks odor of <mark>arsenia</mark>	tric acid, with separation of sulphate of lead.		Jordanite.
1	tor of		In the nitric solution, hydrochloric acid gives a heavy precipitate of AgCl.	POLYBASIT
	rlic o	with hydrochloric acid color		Rionite.
DIVIBION 1	the ga	the flame blue (chloride of copper). A nitric solution is mendered blue by ammonia.	Easily cleavable; the others are not.	ENARGITE
DIV	d give		In the nitric solution, ammonia gives a red- dish-brown precipitate (iron).	Tennantite
	00		Same reaction for iron.	Epigenite.
	han		Gives no precipitate with ammonia.	Binnite.
	~		Gives no sulphur reaction.	DOMEYKIT
	5	Compare Tetrahedrite,	Same.	Algodonite.
	В.	p. 67.	Same.	WHITNEYI
	B.		The concentrated solution is rendered tur- bid by addition of water (bismuth).	Alloclasite.
		Give to the borax bead a sap- phire-blue color.	Gives metallic arsenic in closed tube.	SMALTITE balt-spei
		1	As above.	Skutterudit

* The color of metallic minerals must be observed on a fresh fractor e, as many

CALLIC LUSTRE.

Composition.	Color.*	Streak.	Cleavage or Fracture,	Hard- ness,	Sp. Gr.	Fusibility.	Crystalli- zation.
	Pinkish-white.		Cubical.	1.5—2	8.2-9.7	Easily.	I. (?)
	White.	White.		2.5	10.5	2.—2.5	L
	Yellow.	Yellow.	1	2.5	19.3	2.5-3	I.
.g.	Pale yellow to white,		1	2.5	12.5-15.5	2.5-3	I,
	Copper-red.	Copper-red.		3.	8,9	S .	I.
	Lead-gray.	Lead-gray.		1.5	11.4	1.	I.
Rh,Pd,Fe).	Steel-gray.	Light-gray.		44.5	16—19	Infus.	I.
	Steel-gray.	Steel-gray.		4.5-5.	11.5	Infus.	I.
etc.).	Iron-gray.	Brade.	Octahedral.	4.5	7.5		L
	White,				13.5		L
	Tin-white,	Tin-white.	Granular,	3.5	6.	Vol.	ш.
S ⁵ .	Lead-gray.	Brown.	Basal.	3.	5.5	Easily.	IV.
570		D					
34.	Lead-gray.	Brown,	Basal.	3.	5.3	Easily.	IV.
S*.	Lead-gray.	Black.	Prismatic.			Easily.	IV.
ı) ⁹ (Sb,As)S ⁶ .	Iron-black.	Iron-black.	Uneven.	2.3	6.25	Easily.	IV.
)(As,Bi)?S*.	Iron-black.	Iron-black.					
34.	Iron-black,	Grayish-black.	Prismatic.	3.	4.4		IV.
S1.	Iron-black.	Gray.		8.5-4	4.5	1.5	I.
) ⁵ As ² S ¹² .	Steel-gray.	Black.	Granular.	3.5		Easily.	IV.
S ⁹ .	Steel-gray.	Cherry-red.	Brittle.	4.5	4.4	Easily.	1.
	Tin-white.	Blackish.	Brittle.	3.5		Easily.	Massive.
	Steel-gray.	Bronze.	Tough.	4.		Easily.	Massive.
	Bronze.	Bronze.	Hackly.	8.5	8.3	Easily.	Massive.
,Zn)4(As,Bi)7S*.	Steel-gray.	Black.	Rhombic.	4.5	6.6	Easily.	IV.
Ni)As ^s .	Tin-white.	Gray-black.	Octahedral,	5.5	6.4-7.2	Easily.	I.
	Gray-white.		Cubic.	6.	6.7	Easily.	L

change and become tarnished and dull on exposure to air and light.

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MINERALS WITH METALLIC LUSTRE

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A. Fusible from 1-5, or easily volatile.

Division 1 (continued).

DIVISION &

		General Characters.	Specific Characters.	Species.
			Gives metallic arsenic in the closed tube. The dilute solution gives precipitate with chloride of barium of BaSO ⁴ .	
1		Give to the borax bead a sap-	As above.	Glaucopyrite
		phire-blue color.	Gives no arsenic in closed tube. Dilute so- lution gives a precipitate with chloride of barium of BaSO ⁴ .	
			with cobalt ores, Div. 6, page 69.	
	on charcoal give the parks odor of ar send o		Of a copper-red color.	NICCOLITE (copper- nickel).
	odor		Gives in the closed tube a sublimate of me- tallic arsenic.	Rammelsberg ite.
DIVISION 1.	the park	When dissolved in aqua-regia form an apple-green solution; with ammonia in excess the	rium gives a heavy precipitate.	Gersdorffite (nickel- glance).
E C	sald I	solution becomes sapphire- blue.	Gives a red-brown precipitate with excess of ammonia (iron).	Chathamite (var. Smaltit
	larood		Gives antimony fumes, and a sulphur reac- tion with soda on coal.	Corynite.
	5 2		As above.	Wolfachite.
	8		Compare Ullmannite, Div. 4, p. 67.	
	Ri .	In the closed tube give metallic arsenic, and then fuse, and	Gives sulphur reaction in open tube, soluble in nitric acid, with separation of sulphur; in solution ammonia gives a reddish-brown precipitate (iron).	Arsenopyrit (Mispickel).
		after long heating become magnetic.	Gives only a slight sulphur reaction. In closed tube, after arsenic is driven off, fuses with great difficulty.	Lölingite (Leucopyrite)
		69; Antimony, Div. 4, p. 66; Pyrargyrite, Div. 4, p. 66; Pyrargyrite, Div. 1, p. 72; Geocronite, Div. 4, p. 66; all sometimes containing arsenic		
	D NO.		B. B. volatile without fusion; with soda upon charcoal yields metallic lead.	Lehrbachite.
	the the	With soda in a matrass give metallic mercury.	B. B. fuses and then volatilizes. Gives no lead.	Tiemannite
	an open glass-tube, gtve the wr of selenium.		Gives a reaction for sulphur, in open tube or on charcoal.	Guadalcazar- ite.
		metallic gray, then white, then greenish-yellow subli-	B. B. with soda yields with difficulty lead globules, the nitric solution gives a pre- cipitate with sulphuric acid.	
DIVISION	ed in an tish odor	Give with borax a pure silver globule.	B. B. fuses easily: in O. F., quietly; in B. F., with intumescence.	Naumannite.
	<mark>, or ho</mark> ated in horse-radish o		Solution in nitric acid gives a heavy precip- itate with hydrochloric acid (AgCl).	Eucairite
	charcoal, c	B. B. on charcoal fuse to glob- ules, which after moistening with HCl color the flame azure-blue.	Nitric solution gives a precipitate with sul- phuric acid (PbSO ⁴).	Zorgite.
	B. ON		either sulphune of hydrochloric acid.	Berzelianite.
	4		Contains 18 per cent. of thallium; colors	Crookesite.

METALLIC LUSTRE.

Composition.	Color.	Streak.	Cleavage or Fracture.	Hard- ness,	Sp. Gr.	Fusibility	Orystalling tion.
AsS.	Gray-white.	Black.	Rhombic.	5.	6.	Easily.	IV.
Cu, Sb, As, S.	Gray-white.	Gray-black.		4.5	7.18	Easily.	IV.
	Red-white.	Gray-black.	Cubic.	5.5	6.	Easily.	I.
	Copper-red.	Brown-black.	Uneven.	5.—5.5	7.4	Easily.	 III.
	Tin-white.	Gray-black.	-	5.5	7.	Easily.	IV.
	Gray-white.	Gray-black.		5.5	5.6-6.9	Easily.	I.
Co)As ² .	Gray-white.	Gray-black.	Granular.	· · ·			L
ib)S.	Gray-white.	Black.	Uneven.	4.5-5.	6.	Easily.	L.
3b) S .	Silver-white.	Black.		5.5	6.37	Easily.	<u>IV.</u>
	Silver-white.	Gray-black.	Uneven.	5.5	6.2	2.	IV.
	Silver-white.	Gray-black.		5.5	6.8—8.7	Diff.	IV.
)Se.	Lead-gray.	Black.	Granular.	2.	7.8	Vol.	Massive.
	Lead-gray.	Black.	Granular.	2.5	7.2	Easily.	Massive.
)(8,Se).	Iron-black.	Black.	Compact.	2.	7.15	Easily.	Massive.
	Lead-gray.	Gray-black.	Cubic.	2.5	78.		I.
⁵ Se [‡] ?	Iron-black.	Black.	Cubic.	2.5	8.	Easily.	I.
'Se.	Lead-gray.	Shining.	Granular.			Easily.	Massive.
)Se.	Lead-gray.	Dark-gr ay .	Granular.	2.5	7.5	Easily.	Massive.
	Silver-white.	Shining.		Soft.			Massive.
Ag)Se.	Lead-gray.			2.5	6.9	Easily.	Massive.

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MINERALS WITH METALLIC LUSTRE.

A. Fusible from 1—5, or easily volatile.

DIVISION 8.

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DIVERSION & (in part).

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1. MINERALS WIT

nteh- hya- gray	General Characters,	Specific Characters,	Species.
olors the R. F. green, or greents. rated sulpharic acid, give a hyd- diagpears, yleiding a black-gray	Wholly volatile; fuses easily; fumes strongly.	Burns with a greenish flame. Soluble in nitric acid.	Tellurium.
F. grewrio ac	Wholly soluble in nitric acid.	The solution gives a heavy precip. with sul- phuric acid. Soft but not malleable.	Altaite.
the R.		B. B. with soda gives globule of silver; mal- leable.	Hessite.
ting which colors of a concentrated on of water disap	Soluble in nitric acid with the separation of gold.		Müllerit (sylvanite o taining les
tting tolis of the conco	On charcoal, with sulphur and iodide of potassium, gives a red sublimate (bismuth).	Fuses easily to a brittle silver-white globule.	TETRAD MITE.
ed u ditt		Gives after a little blowing the selenium odor.	Joseite.
s a tohite tily heat ch on ad tum.	After long heating gives a malle- able metallic globule. Incom- pletely soluble in nitric acid.	Soluble in aqua-regia, with separation of chloride of silver.	Sylvanite (graphic te rium).
gie volu		Same reactions (contains more silver).	Petzite.
B. On charcoal give a white control which colors blue. In a two genity hadda with concentrated onth-red color. which on addition of water diago precipitate of latitutian.	Heated with strong sulphuric acid gives a hyacinth-red or brownish - yellow solution; not a pure red like preceding.	Soluble in aqua-regia; in the solution SO ₃ gives a precipitate (lead).	Nagyagit (foliated tellurium
B. B. diud	Compare Aikinite, Div. 5, p. 68.		
- Auo		Tin-white. B. B. takes fire and continues to burn without further heating, and becomes covered with white needles of oxide of antimony.	
ue antim	S potassa is rapid for the most p Gives in nitric ac blue color; this a white precipi is rendered vio	When pulverized and treated with causti potassa is rapidly colored ochre-yellow, and for the most part dissolved.	
stube, give den		Gives in nitric acid a partial solution of a sky- blue color; this, with sulphuric acid, yields a white precipitate of sulphate of lead, and is rendered violet-blue on addition of an excess of ammonia.	BOURNONI
glas	B. B. are nearly or completely	Same reaction, but the aqua-regia solution is not precipitated by sulphuric acid.	Stylotypite
I AVISION The open flo	volatile in a continued blast.	Oxidized by nitric acid to a white powder, imparting no color to the acid.	Jamesonit
an -		Same as Jamesonite, not cleavable.	Zinkenite.
al, 0		As above.	Boulangeri
Auros		As above (sometimes contains arsenic).	GEOCRONI
on c		As above.	Plagionite.
RÌ		As above.	Meneghini
8		Fused with sulphur and iodide of potassium gives a bismuth reaction.	Kobellite.
		Compare Galenite, Div. 5, p. 67.	

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FALLIC LUSTRE.

Composition.	Color.	Streak.	Cleavage and Fracture.	Hard- ness.	Sp. Gr.	Fusibility.	Orystalling tion.
- <u></u>	Tin-white.	Tin-white.	Hexagonal.	2.—2.5	6.2	1.	ш.
	Tin-white.	Tin-white.	Cubio.	83.5	8.1	1.	I.
).	Lead-gray.	Gray.	Sectile.	2.5—3.	8.5	1.	IV.
3,Pb,Te, Sb .	Brass-yellow.			2.	8.	Easily.	v .
8.	Steel-gray.		Basal.	1.5—2.	7.5	Easily.	ш
e ⁴ SeS ³ .	Black-gray.		Basal.	1.5-2.	7.9	Easily.	ĪĪĪ.
u)Te².	Steel-gray.	Steel-gray.	Prismatic.	1.5—2.	8.	Easily.	IV.
u) ² Te.	Iron-black.	Iron-black.		2.5	9.	Easily.	IV.
u)(S,Te) ² .	Black-gray.	Gray.	Basal-fol.	1.—1.5	7.	Easily.	п.
	Tin-white.	Tin-white.	Perfect.	8.5	6.9	1.	 III.
	Lead to steel- gray.	Lead-gray.	Prismatic.	8 .	4.5	1.	1▼.
5b8 ° .	Steel-gray.	Iron-black.		2.5—8.	5.8	1.	I V.
g ³ ,Fe) ³ Sb ² S ⁶).	Iron-black.	Iron-black.	-	8.	4.7	1.	IV.
²8⁵.	Lead-gray.		Basal.	23.	5.6	1.	I V.
S4.	Lead-gray.		-	8.—8.5	5.8	1.	IV.
[*] 8•.	Lead-gray.			2.5—8.	5.9	1.	IV.
²Sª.	Lead-gray.		Granular.	28.	6.5	1.	IV.
6g18.	Lead-gray.			2.5.	5.4	1.	v.
'S'.	Lead-gray.			2.5	6.8	1.	v .
ShS ⁴ .	Lead-gray.			Soft.	6.8	1.	Fibrous.

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MINERALS WITH METALLIC LUSTRE.

A. Fusible from 1-5, or easily volatile.

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DIVERSON 4 (continued).

DEVISION 5 (in part).

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1. MINERALS WITE

	General Characters,	Specific Characters.	Species.
		Gives no sulphur reaction.	Dyscrasite (antimonial silver).
Drvisios 4.—(Continued.) or in the open glass-tube, gles dense antimony fumes.	B. B. give with a mixture of borax and soda a malleable silver-bead, and the nitric solution yields with hydro- chloric acid a precipitate of	a violet-blue color (copper).	FREIBERGITE (silver tetrahe drite).
e anti	chloride of silver.	Gives sulphur reaction, but no blue with ammonia.	STEPHANITE.
ense		As above.	Miargyrite.
('Deur		As above ; the nitric solution gives a precip- itate of sulphate of lead with H ³ SO ⁴ .	Brogniardite.
utto ibe,		Same.	Freieslebenite
+ (10		Compare Pyrargyrite, Div. 1. p. 72.	1.200
Drviston 4.—(Continued.) 1 the open glass-tube, glos (Heated in a glass tube with soda gives a sublimate of mercury.	m	SPANIOLITE.
Drv1 or in the	si Give after long heating on charcoal a magnetic glob- Easily d	The nitric solution gives no precipitate with hydrochloric acid, but usually gives reac- tions for iron and zinc.	
harcoal,		Very closely resembling the above in blow- pipe reactions is the rare	Chalcostibite (antimonial- copper).
-		Gives no sulphur reaction in open tube ; dif- ficultly fusible; but little acted on by hy- drochloric acid; completely dissolved by aqua-regia.	n the second
		Easily dissolved in hydrochloric acid, with disengagement of sulphuretted hydrogen.	BERTHIERITE
		Easily fusible; hydrochloric acid has little effect; aqua-regia dissolves it with sepa- ration of sulphur.	
the open eddens a d, but do	Malleable, can be cut with a knife like lead. In the nitric	The nitric solution is colored blue by excess of ammonia ; moistened with HCl colors the flame blue.	
sced in which r the en	solution hydrochloric acid Does not give the above reacting gives a heavy precipitate of soda gives a globule of silver.		ARGENTITE (silver-glance).
, or ple ction, a aced in divisio	chloride of sliver.	Differs only in crystalline form from argen- tite.	Acanthite.
rea rea	The roasted minerals give with	The powder is leek-green.	Alabandite.
on 5. read	(manganese).	The powder is brownish-red.	Hauerite.
DIVIEION e a sulphur rea sulphurous acie blue litmus pay ons of the prece	Streak red; mixed with soda in	Many varieties have a gray to black color, but the streak is red. The rare <i>Metacinnabarits</i> is amorphous HgS; has a black streak, H=3; G=7.72.	Cinnabar.
D. B. B. with soda give a sulphu glass-tube give a sulphu sirip of moistened blue it not give the reactions of	B. B. with soda on coal gives a lead-globule and covers the coal with a yellow coat (oxide		Galenite (ga- lena).
B. B. w glass- surip not gi	of lead).	Cuproplumbite and Huascolite are re- spectively cupriferous and zinciferous varieties of galena.	

FALLIC LUSTRE.

	1					
Silver-white.	Silver-white.	Basal.	8.5-4.	9.6	1.5	IV.
Steel-gray.	Gray.		3.5	4.8		L
Iron-black.	Black.	-	2.5	6.26	1.	īv.
Iron-black.	Cherry-red.		2.5	5.2	1.	v .
Black-gray.			8.	5.9	Easily.	L
Steel-gray.	Gray.	Prismatic.	22.5	6.—6.4	Easily.	<u>v.</u>
Iron-black.	Red-brown.	-	8.5	5.1	1.5	I.
Lead-gray.	Dark-gray.		3.5	4.5-5.1	1.5	I.
Lead-gray.	Black.	Prismatic.	3.5	4.8	1.	I V .
Copper-red.	Red-brown.		5.5	7.5	8.	ш
Steel-gray.			2.—3,	4.8	1.5	IV. ?
Steel-gray.			5.5	6.8	8.	L
Gray-black.	Gray-black.	Malleable.	2.5	6.8	Easily.	I.
Gray-black.	Gray-black.	Malleable.	2.5	7.9	1.5	l.
Gray-black.	Gray-black.	Malleable.	2.5	7.2	1.5 .	IV.
Iron-black.	Green.	Cubic.	8.5	4.	8.	I.
Brown-black.	Brown-red.		4.	8.46	8.	I.
Red.	Red.	Perfect.	2.5	8.9	Vol.	ш.
Lead-gray.	Gray.	Cubic.	2.5	7.5	1.	L
	Steel-gray. Iron-black. Black-gray. Steel-gray. Iron-black. Lead-gray. Copper-red. Steel-gray. Steel-gray. Gray-black. Gray-black. Brown-black. Brown-black.	Steel-gray. Gray. Iron-black. Black. Iron-black. Cherry-red. Black-gray. Gray. Steel-gray. Gray. Iron-black. Bed-brown. Lead-gray. Dark-gray. Lead-gray. Black. Copper-red. Bed-brown. Steel-gray. Black. Gray-black. Gray-black. Gray-black. Gray-black. Gray-black. Gray-black. Iron-black. Gray-black. Brown-black. Gray-black. Brown-black. Brown-red. Brown-black. Brown-red.	Steel-gray. Gray. Iron-black. Black. Iron-black. Cherry-red. Black-gray. Gray. Steel-gray. Gray. Iron-black. Red-brown. Iron-black. Bed-brown. Lead-gray. Dark-gray. Lead-gray. Black. Prismatic. Copper-red. Red-brown. Steel-gray. Black. Steel-gray. Black. Steel-gray. Black. Gray-black. Gray-black. Gray-black. Gray-black. Malleable. Gray-black. Gray-black. Gray-black. Brown-black. Green. Brown-black. Brown-red. Brown-black. Brown-red.	Steel-gray. Gray. 3.5 Iron-black. Black. 2.5 Iron-black. Cherry-red. 2.5 Black-gray. Gray. 8. Steel-gray. Gray. Prismatio. Iron-black. Bed-brown. 3.5 Iron-black. Bed-brown. 3.5 Lead-gray. Dark-gray. 3.5 Lead-gray. Black. Prismatio. Copper-red. Red-brown. 5.5 Steel-gray. 3.5 3.5 Gray-black. Gray-black. Prismatio. Steel-gray. 5.5 5.5 Gray-black. Gray-black. Malleable. Gray-black. Gray-black. Malleable. Gray-black. Gray-black. Malleable. Iron-black. Gray-black. Stown-red. Hed. Brown-red. 4. Red. Red. Perfect. 2.5	Steel-gray. Gray. 3.5 4.8 Iron-black. Black. 2.5 6.26 Iron-black. Cherry-red. 2.5 5.2 Black-gray. Gray. Prismatio. 22.5 66.4 Iron-black. Red-brown. 8.5 5.1 Lead-gray. Dark-gray. 8.5 5.1 Lead-gray. Dark-gray. 8.5 4.8 Copper-red. Bed-brown. 8.5 7.5 Steel-gray. Black. Prismatio. 3.5 4.8 Copper-red. Bed-brown. 5.5 6.3 7.5 Steel-gray. Image: Steel-gray. 1mage: Steel	Steel-gray. Gray. 3.5 4.8 Iron-black. Black. 2.5 6.26 1. Iron-black. Cherry-red. 2.5 5.2 1. Black-gray. Gray. Prismatio. 22.5 66.4 Easily. Steel-gray. Gray. Prismatio. 22.5 66.4 Easily. Iron-black. Red-brown. 3.5 4.5-5.1 1.5 Lead-gray. Dark-gray. 3.5 4.5-5.1 1.5 Lead-gray. Black. Prismatio. 3.5 4.8 1. Copper-red. Red-brown. 5.5 7.5 8. Steel-gray. Black. Prismatio. 3.5 4.8 1.5 Steel-gray. Black. Prismatio. 3.5 4.8 1.5 Gray-black. Gray-black. Malleable. 2.5 6.8 Easily. Gray-black. Gray-black. Malleable. 2.5 7.3 1.5 Iron-black. Green. Cubic. 8.5 4. 8.6 Brown-black. Br

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MINERALS WITH METALLIC LUSTRE.

A. Fusible from 1-5, or easily vointile.

Devianow 5 (montinued).

I. MINERALS W

		(Jeneral Characters.	Specific Characters.	Spe
		pper. n of	With sulphur and io- dide of potassium	Gives with soda on coal a globule of copper.	Wittich (cup) bism
	40	of col		In the nitric solution sulphuric acid gives a precipitate of PbSO ⁴ .	Aikinit
	ptil so	nide on a	dide of bismuth. The saturated nitric	Gives with soda on coal a strongly magnetic globule containing nickel.	Grunau
		din	solution is rendered turbid by addition	Same as Aikinite.	Chiviat
		24	of water,	Same as Wittichenite.	Empleo
	in the end	color o on, whi ne.		Brass-yellow color.	Chalco (copp rites)
	ed	blue	1	Brass-yellow color; cleavage cubic.	Cubani
	ons.	the bl n sol	Dura to brittle steel	Brass-yellow color; the fresh fracture tar- nishes to a golden-yellow in 24 hours.	Barnha
	e titmus pap	Moistened with HCl give to the flame the blue color of chloride of copper. Give with nitric acid a blue or green solution, which upon addition of ammonia in excess becomes violet or aky-blue.	Fuse to brittle steel- gray magnetic glob- ules,	Color bronze-yellow, exposed surfaces pur- ple, whence it is called variegated copper; in the nitric solution sulphuric acid gives no precipitate.	Bornite egate per).
	ened blu f the pre	dive to id a bl		Resembles bornite, but in the nitric solution sulphuric acid gives a precipitate of sul- phate of lead.	Castilli
('pen	which reactions a strip of motstened blue titmus paper placed in the end ; or with soda but do not give the reactions of the preceding divisions.	HOI E	Give none of the above reactions.	The nitric solution gives with HCl a heavy precipitate of AgCl.	Strome
DIVISION 5 (Continued.		d with ith nita		B. B. in O. F. alone on coal yields a globule of copper. Soluble in nitric acid with separation of sulphur.	Chalo (cop glan
IBION D		B. B. partially reduc		Gives by itself no metallic malleable globule. Soluble in nitric acid, with separation of sulphur and binoxide of tin.	STAN (tin py
AI				Compare Tetrahedrite, Div. 4, p. 67.	-
H	acid, whit		B. B. partially reduced to silver; the partial nitric solution gives a heavy precipitate of chloride of silver with HCl.		
	uphurous phur rea			The roasted mineral gives to the borax bead a sapphire-blue color. Soluble in nitric acid, forming a rose-red solution.	
	the all and	ed with hydro color to the fi	Gives like reactions, but also, when moisten- ed with hydrochloric acid, imparts a blue color to the flame.	Carrolli	
	in the open glass-tube f		the might color while hat noddish brown	(canilla	
	u.			As above, but gives sulphur in the closed tube.	Beyrich
	the op			Magnetic before fusion; color pinchbeck- brown. Reacts for nickel.	Pentlar
	Hoted in			Give only the reaction for iron. Magnetic before fusion, gives but little sulphur in the closed tube.	Pyrrl (magne rite
				Gives only the reactions of iron; not magne- tic before fusion. Gives sulphur in the closed tube.	Pyrite pyri
		Compar	e Sphalerite.	Same as for pyrite ; can be distinguished only by crystalline form.	Marc (white i

TALLIC LUSTRE.

Composition.	Color.	Streak.	Cleavage or Fracture.	Mard- ness.	Sp. Gr.	Fusibilit y .	Crystalline tion.
i8³.	Steel-gray.	Black.		3.5	4.6—5.	Easily.	IV.
BiS ³ .	Lead-gray.		-	22.5	6.7	1.	IV.
,Cu,Fe,S.	Steel-gray.	Dark-gray.	Octahedral.	4.5	5.1	Éasily.	jr.
"u2)2Bi6S11.	Lead-gray.		-	-	6.92	1.	Massive.
52.	Tin-white.			-		Easily.	IV.
S³.	Brass-yellow.	Green-black.	Uneven.	8.5	4.8	2.	п.
² S ⁴ .	Bronze-yellow.	Red-bronze.	Cubic.	4.	4.1	Easily.	I.
e²S⁵.	Bronze-yellow.		-	8.5	4.5	Easily.	Massive.
uFe)S.	Bronze-yellow, purple.	Black.		8.	5.	Easily.	I.
b,Fe,Zn,Ag,S.	Copper-red.	Black.	Foliated.	8.	5.2	Easily.	Massive.
['] u) ² S.	Steel-gray.	Gray.		2.5-3.	6.8	Easily.	IV.
	Steel-gray.	Gray.	Conchoidal.	2.5-3.	5.7	Easily.	IV.
n,Fe,Zn)S.	Steel-gray.	Black.		4.	4.4	Easily.	Massive.
²8³ .	Pinchbeck- brown.	Black.	Basal.	1.—1.5	4.2	Easily.	1V.
ï) ⁸ S ⁴ .	Steel gray.	Black-gray.		5.5	4.9	Easily.	I.
18.	Tin-white.	Gray.	•	5.5	4.85	Easily.	I.
	Bronze-yellow	Bright.		8.—8.5	5.6	Easily.	III.
	Lead-gray.			33.5	4.7	Easily.	<u>III. ?</u>
i) 8.	Bronze-yellow.	Brown.		8.5-4.	4.6	Easily.	I
	Bronze-yellow	Black-gray.	Basal.	4.	4.5	Easily.	ш.
······	Brass-yellow.	Brown-black.		6.—6.5	4.9	Easily.	1.
	Pale-yellow to white.	Black-gray.		6.—6.5	4.7	Easily	IV.

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MINERALS WITH METALLIC LUSTRE.

A. Fusible from 1-5, or easily volatile.

DIVISION 5 (concluded).

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DEVISION G.

L MINERALS WITH

tinued.) age.	General Charaotera.	Specific Characters.	Species.
DAVISION 5 (Continued. See preceding page.	With sulphur and iodide of po- tassium give, on coal, a red	B. B. fuses in R. F. with effervescence, giv- ing a globule of bismuth, and a bismuth coating on coal. Soluble in nitric acid.	ITE (bismuth
See p	sublimate of iodide of bis- muth.	As above, but gives a precipitate of sulphate of lead with sulphuric acid.	Chiviatite.
A		S Compare Bismuth, below.	
	B. B. in a matrass yield me- tallic mercury and leave a		Amalgam.
	spongy mass of silver.	Yields less mercury in the closed tube.	Arquerite.
	Fused with sulphur and iodide of potassium, coats the coal with a red sublimate of iodide of bismuth.	In the open glass-tube gives almost no fumes, and the metal becomes surrounded with fused oxide, which is dark-brown when hot, and yellow when cold.	BISMUTH,
	Colors the borax bead cobalt- blue.	Heated with phosphoric acid gives a violet solution (manganese).	Rabdionite.
	Difficultly fusible. Heated in B. F. becomes magnetic.	Streak, cherry-red.	Hematite (specular iron
	With soda on charcoal easily reduced to metallic copper.	Compare Cuprite and Melaconite, Div. 3, p. 75, sometimes with metallic lustre.	
stons.	Magnetic before heating.	Generally fusible above 5.	Magnetite (magnetic iron).
ditte		Gelatinize with hydrochloric acid. Some-	HORTONOLITE,
Dup		times magnetic from associated magnetite.	FAYALITE.
DIVISION 6. Not belonging to the foregoing divisions.	The fine powder, boiled with aqua-regia, gradually assumes a yellowish color.		
Not belong	With borax in O. F. give an amethystine bead.	Compare Rhodonite, Div. 5, p. 87, sometimes altered to a black metallic hy- drous silicate; Klipsteinite, Div. 4, p. 85; and Psilomelane, Div. 1, p. 70, which in some varieties is fusible.	
	Gelatinize perfectly with hydro-	Easily fusible, swells up but slightly. (See Div. 5, p. 78.)	ILVAITE.
	chloric acid.	Easily fusible, swells up strongly. (See Div. 5, p. 78.)	ALLANITE.
	With soda easily reduced to metallic lead.		Plattnerite.
	obtained, which neutralized If the latter is digested with	d with water and filtered, a green solution is with HCl gives a light-colored precipitate. strong HCl, and boiled with tin, then diluted it gives a clear sapphire-blue solution.	a

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CALLIC LUSTRE.

Composition	Color.	Streak.	Cleavage or Fractors.	Hard- ness.	Sp. Gr.	Fusibility.	Crystallin tion.
<u></u>	Lead-gray.	Gray.	Prismatic.	2.	6.4	1.	IV.
S ¹¹ .	Lead-gray.	Gray.	Foliated.		6.92	1.	
and Ag ² Hg ³ .	Silver-white.	Gray.		 3.	13.7—14	1.	 L.
ζ.	Silver-white.	Gray.	Malleable.		10.8		I.
	Reddish-white	White.	Basal.	2.5	9.7	1.	ш
n,Co)(Mn,Fe)O4.	Black.	Metallic greasy streak.		1.	2.8	3.	Stalact
	Steel-gray to black.	Red.	Scaly, fibrous, compact.	5.5 <u>-</u> 6.5	5.	Infus.	ш.
<u> </u>	Iron-black.	Black.	Octahedral.	 5.56.5	4.9—5.2	5.	L.
g) ² SiO ⁴ .	Yellow-black.	Dirty-white.	Prismatio.	6.5	3 .9	4.	IV.
)4.	Black.	Brown.	Prismatio.	6.5	4.1	8.	IV
n)WO4.	Black.	Black.	Prismatic.	5. 5	7.8	8.	₹.
Fe ⁴ FeSi ⁴ O ¹⁸ .	Black.	Black.		5.5-6.	3.8-4 .	2.5	IV.
.,Di,Fe,Ca) ³ (Al, 5i ² O ¹³ .	Brown-black.	Gray.		5.5-6.	8.—4.2	2.5	٧.
	Iron-black.	Brown.			9.8		
,UO ³) ⁶ (Cb,Ta)4O ⁴	Velvet-black.	Dark red- brown.		5.5-6.	5. 6	4.5	IV.

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MINERALS WITH METALLIC LUSTRE.

3. Infusible or fusible above 5, and non-volatile.

DIVISION 1.

DEVICION &

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L MINERALS W.

Sonas-	General Characters.	Specific Characters.		
oes to the		Moistened with HCl colors the outer flame beautifully blue (chloride of copper).		
DIVISION 1. In O. F. a very small quantity gives to the boad the amelihysithe red of manganese.	With hydrochloric acid evolve chlorine. Contain little or no water.	Color brownish-black.		
DIVISION 1. mail quan		Color iron-black to steel-gray.		
thery		Prismatic cleavage very perfect.		
bead the d	Yield much water in the closed tube.	In the hydrochloric solution sulphuric acid generally yields a white precipitate of sul phate of baryta.		
B. B. 4	Div. 2, below. Hauerite and Alabandite, Div. 5, p. 67.			
	Streak red.	Decrepitates, and gives much water in th closed tube.		
DIVISION 2. on charcoal after iong heating in R. F. become so.		Slowly soluble in hydrochloric acid.		
	Magnetic without heating (sometimes but slightly).	With soda gives the manganese reaction and on coal in R. F. gives a faint yello sublimate (ZnO).		
		 Strongly magnetic, does not give above ractions. Difficultly fusible. In the solution after the oxidation of t protoxide of iron with chlorate of pota and its precipitation with an excess of armonia, phosphate of soda gives a precipitate of the ammonio-phosphate of magnetic gives a stronmanganese reaction. 		
che		Compare Menaccanite, below.		
B. B.	soda, dissolved in HCl., gives a solution which, boiled with	phate of potash.		
or	tin-foil, gradually assumes a violet color.	125" Rutile, Anatase and Arkansite some times become magnetic after long heating		
Are magnetic,	Streak ochre-yellow (some- times has a sub-metallic lustre).	Much water in the closed tube.		
4	Div. 4, p. 92, sometimes with metallic lustre; also the min- erals of the following section, especially <i>Chromite</i> .			

ETALLIC LUSTRE.

	Composition.	Color.	Streak.	Oleavage or Fracture.	Hard- ness,	Sp. Gr.	Orystalli- sation.
е.	Cu ³ Mn ² O ⁹ .	Black.	Black.	Basal.	4.5	5.	v
E.	3MnO ³ +MnSiO ³ .	Brown-black,	Black.		66.5	4.7	- II.
nite.	Mn ² O ⁴ .	Brown-black.	Chestnut- brown.	Basal.	55.5	4.7	п.
ю.	MnO ³ .	Iron-black.	Black.		2.—5.	4.82	IV.
TE.	H ^a MnO ⁴ .	Steel-gray.	Red-brown.	Prismatic.	4.	4.8	IV.
ANE	(Mn,Ba ₉ K ³) ⁵ O ⁹ +aq.	Black.	Brown-black.		5.—6.	8.7—4.7	Amorph.
(hy- ;ite).		Beddish-black	Red.	Fibrous, compact.	56.	4.14	_
ite iron)	₽eO³.	Reddish-black	Red.	Scaly, fibrous, compact.	66.5	4.5-5.8	ш.
ite.	(Fe,Zn,Mn)(Fe,Mn)O4.	Iron-black.	Reddish- brown.		5.5-6.5	5.1	I.
e.	Fe ³ O ⁴ .	Iron-black.	Black.	Octahedral.	5.56.5	4.9-5.2	I.
	(Mn,Mg)(Fe, M n)O ⁴ .	Black.	Black-brown.		6.	4.75	I.
	Mg∓eO4.	Black.	Black.		6.—6.5	4.5-4.6	L.
nite ron).	(Ti,Fe) ² O ³ .	Black.	Black.		5.—6.	4.55.	III.
te ema-	 Н'Fe'O'.	Brown.	Yellow.	Fibrous, compact, earthy.	5.—5.5	8.6-4.	

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MINERALS WITH METALLIC LUSTRE.

B. Infusible or fusible above 5, and non-volatile.

Devesion 1.

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L MINERALS W

			General Characters.	Bpecião Characters.				
			Imparts a beautiful emerald- green color to the beads of borax and salt of phospho- rus when cold.	Sometimes strongly magnetic; only slightly attacked by hydrochloric acid.				
			Which with soda on coal is reduced to metallic tin.					
volatile			Very soft; soil the fingers.	B. B. in forceps colors the flame light-green on charcoal with soda gives a sulphur re action, and gives coating of molybdic acid				
d non-						Does not give the above, but deflagrate: with nitre, affording carbonate of potassa		
e 5, an		egoling d	Give to the salt of phosphorus- bead the violet color of titanic acid.	Orystallizes in cubes.				
Aoqu	DIVINION 8.	the for		Compare Rutile and Brookite, Div. 6. p. 95, sometimes with metallic lustre.				
usible	DIAT	DIVISION 3. Not included in the foregoing divisions.	nded in I	huded in I	inded in 1	luded in	Fused in a matrace with nitre evolves the peculiar odor of oxide of osmium.	Not perceptibly attacked by borax, salt of phosphorus, or nitro-hydrochloric acid.
le or f				B. B. immediately changes its color to yel- low or white.				
B.—Infusible or fusible above 5, and non-volatile			Slightly attacked by acids.	The powder fused with bi-sulphate of pot- ash, then boiled with HCl, and filtered and the liquid evaporated with addition of tin-foil, it assumes a beautiful blue color, which rapidly fades, and gradually disappears upon addition of water.				
				Gives like reactions.				
				Compare Polyerase, Div. 4, p. 93 Allechynste, Div. 6, p. 95.				
			a yellow fluid, from which	With salt of phosphorus in R. F. a greer bead, becoming yellow in O. F. Evaporated with phosphoric acid gives ar emerald-green solution.				

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ETALLIC LUSTRE.

L	Composition.	Color,	Streak,	Cleavage or Fracture.	Hard- ness.	Sp. Gr.	Crystalling tion,
ite iron)	FeGrO4.	Iron-black.	Brown.	Uneven.	5.5	4. 3	L
nite.	MoS [‡] .	Blue-gray.	Greenish.	Foliated.	1.—1.5	4.6	V. ?
•	Ο.	Iron-black.	Black.	Foliated.	1.—2.	2.	щ
э.	CaTiO ⁴ .	Iron - black to yellow.	Gray.		5.5	4.08	I.
ne.	Ir,Os,Rh,Ru.	Tin-white.	Gray.		6.—7.	19.8-21.1	ш.
alite	(Fe,Ca,Y) ² (Ta,Cb) ² O ⁷ .	Yellow to black.	Grayish.	Conchoidal.	5.5	5.7	IV.
	Fe(Mn)Ta ² O ⁴ .	Black.	Brown-black.	Brittle.	6.—6.5	7.—8.	I V.
т е .	FeCb ² (Ta ²)O ⁶ .	Black.	Red-black,	Brittle.	6.	5. 4 —6.5	IV.
ite.	(Y,E,Ce,Fe) ³ (Cb,Ta) ² O ⁸	Black.	Pale-brown.	Conchoidal.	5.5—6.	5.8	п.
nte nde).	U'0'.	Brownish- black.	Brown-black.	Conchoidal, uneven.	5.5	6.4-7.	L

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MINERALS WITHOUT METALLIC LUSTRE

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A. Easily volatile or combustible.

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B. Fusible from 1-5, and non-volatile.

L Yield a metal or a magnetic mass with soda.

Orvince L

		General Characters,	Specific Characters,	Spe
		B. B. burns with a blue flame, emitting the odor of sulphu- rous acid.		Sulphu
1.7		Fuse readily and volatilize, and	Color aurora-red.	Realga
ble.		B. B. on coal with soda in R. F. give off arsenical fumes.	Color lemon-yellow.	Orpim
busti		Heated in a closed tube give a crystalline sublimate. B. B.	Occurs in thin plates with perfect cleavage.	Claude
COL		with soda on coal give the strong garlic odor.	Color white.	Arsenol
AB. B. easily volatile or combustible.		Easily fusible and volatile, cov-	Dissolve mostly in HCl with the evolution of sulphuretted hydrogen. Heated with pot- ash solution the powder becomes yellow.	KERME
vola		ering the coal with a white sublimate (oxide of anti- mony). Insoluble in water.	gas. Unchanged by potash.	VALEN
asily		mony). Insolutio in water.	Same reactions as Valentinite; differs only in crystalline form.	SENAL TIT
B, e		Volatilize with dense white fumes; soluble in water.	Volatile without fusion, its solution gives no precipitate with chloride of barium.	SAL A
A-B.		Treated with potash solu- tion give an ammoniacal odor.	Volatile with fusion, its solution gives a heavy precipitate with chloride of barium (sulphate of baryta).	Masong
-		With soda in the closed glass-	Streak red. Gives a reaction for sulphurous acid in the open glass-tube.	Cinnah
		tube give a sublimate of me- tallic mercury.	Streak white. In the nitric solution nitrate of silver gives a heavy precipitate of the chloride of silver.	Calome
		Partly volatile; with soda on coal gives lead globules.	Deposits a lead-coating on charcoal.	Cotunn
		See also mineral coals in the appendix.		1
globule or			B. B. on coal gives an arsenical odor.	(light- ver or
etallic gl	ute of silver.	Streak red.	B. B. on coal gives a white sublimate of oxide of antimony.	Pyran (d a r silver
give a m mass.	DIVISION 1. charcoal give a globule of		Same reactions as Proustite, but easily dis- tinguished by its orange-yellow color and streak.	Xantho
tic	NO		Compare Miargyrite, Div. 4, page 67.	
soda on charcoal give a metallic a magnetic mass.	DIVISION Boda on churcoal g		In a closed tube, with bisulphate of potassa, gives off hydrochloric acid vapors, fuses to a pale hyacinth-red globule, becomes yel- low when cold.	CERAR ITE.
with soda	totth	Malleable and sectile.	In closed tube, with bisulphate of potassa, gives off iodine vapors, fuses to a very dark, almost black globule.	Iodyrite
B. wi	B.B.		In a closed tube, with bisulphate of potassa, gives off bromine vapors, fuses to an in- tense garnet-red globule, becoming yellow when cold	

In minerals without metallic lustre there is frequently a wide range of color in a single species (for example, in Tou connection with other characters in determining non-metallic mineral species. The streak in this group is generally paler t + If a compound of iodine is fused with sulphide of bismuth, a red coat is formed on the coal or in the open glass tut applicable to the haloid salts of silver. The bismuth sulphide is made by fusing metallic bismuth with sulphur, and grir

LETALLIC LUSTRE.*

Composition.	Color.	Cleavage or Fracture,	Lustre.	Hard- ness,	Sp. Gr.	Fusibility.	Crystalling tion.
	Salphur - yel- low.	Conchoidal.	Resinous.	1.5-2.5	2.	Easily.	17.
	Aurora-red.	Conchoidal.	Resinous.	1.5-2.	3.5	Easily.	v .
	Lemon-yellow	Foliated.	Pearly.	1.5-2.	3.48	Easily.	I V.
	White.	Prismatic.	Pearly.	2.5	3.8		I V .
	White.		Silky.	1.5	3.69	Vol.	I.
).	Cherry-red.	Basal.	Adamantine.	1.5	4.5	1.	v.
	White.	Prismatic.	Adamantine.	2.5-8.	5.56	I .	IV.
	White.	Octahedral.	Adamantine.	2.—2.5	5.22		L
l.	White, yellow.		Vitreous.	1.5-2.	1.53	Vol.	I.
$^{2}SO^{4} + aq.$	White, gray, yellow.		Vitreous.	2.	1.7	Vol.	IV.
	Red.	Hexagonal.	Adamantine.	2.—2.5	9.	Vol.	Ш.
	Gray-white.		Adamantine.	1.5	6.5	Vol.	п.
	Yellow-white.		Adamantine.	2.	5.2	Easily.	IV.
S³.	Cochineal-red.	Conchoidal.	Adamantine.	2.5	5. 8	1.	 III.
83.	Dark - red to black.	Conchoidal.	Adamantine.	2.5	5.8	1.	1 II .
⁸ S ¹⁰ .	Pomegranate- yellow.		Adamantine.	2.	5.1		ш.
	Pearl-gray.		Resinous ada- mantine.	1.—1.5	5.5	I .	 L.
	Lemon-yellow	Basal.	Adamantine.	1.5	5.7	I.	ш.
	Greenish-yel- low.		Adamantine.	28.	5.8 —6 .	I .	L.
,Br).	Green to dark- yellow.		Adamantine.	11.5	5.3-5.8	I.	I.

sy other species, it varies from colorless to black); color, can therefore, generally be used only as confirmatory in mineral, and in a large majority of non-metallic minerals it is very nearly where, compound similarly treated gives a yellow cost, and a chlorine compound a white cost. These reactions are especially π to a powder. (V. Goldschmidt.)

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MINERALS WITHOUT METALLIC LUSTRE

B. Fusible from 1-5, and non-volatile.

L Yield a metal or a magnetic mass with sods

DIVISION \$ (in part).

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II. MINERALS WITH

			General Characters.	Specific Characters.	Species.
			B. B. on charcoal give coatings of lead and antimony.	Fused in a salt of phosphorus bead which has been saturated with oxide of copper, colors the flame blue (chloride of copper).	Nadorite. 33
	1			Gives water in the closed tube.	BINDHEIMITA.
				Fused in forceps in R.F. crystallizes on cool- ing. (Like Pyromorphite.)	MIMRTITE, 198
			B. B. on charcoal give arsen- ical odors.	A variety of Mimetite containing phosphate of lime; gives the reaction for phosphoric acid.	Hedyphane.
olatile.	a magnetio ma u .		The cold nitric solution gives with molybdate of ammonia a yellow precipitate (phos- phomolybdate of ammonia).	plainly crystalline; gives the chlorine reac- tion with sell of nhosphorus and oxide of	ite
ially v	ubout 1			Imparts to the borax bead an emerald-green color, which in O.F. becomes light olive- green, then yellow, and finally colorless.	Dechenite (aræoxene).
dy part	8	of lead.	Color red.	Imparts to the borax bead an emerald-green color, which is constant in both flames. Streak orange.	CROCOITE (chromate off lead).
	a metallic globule	a alobule o		As above. Streak brick-red.	Phœnico- chroite.
ELIG,	etalli			Gives with borax a yellow glass, which be- comes colorless on cooling.	Minium.
Fusible from 15, and not volatile, or only partially volatile.	m a sa	DIVISION 2. charcoal give		With soda gives the reaction for sulphur. Heated with nitric acid sulphate of lead separates. Gives water in closed tube.	Linarite.
ad ne	ng po	DIVI on chai		The solution gives with nitrate of silver a precipitate of AgCl.	Phosgenite.
	eharcoal give	soda		The partial solution gives with nitrate of baryta a precipitate of BaSO ⁴ .	Lanarkite.
OB	da on	B. with	Dissolve in nitric acid with effervescence.	Not affected by the above reagents.	Cerussite (white lead).
ble fr	with soda	ä.		The same as lanarkite, but is orthorhombic in crystallization.	Leadhillite.
F usi	B. w			The same as lanarkite, but is hexagonal (rhombohedral).	Susannite.
	m			Prismatic cleavage very perfect.	Mendipite.
на .	H		effervescence. The solutions give heavy preci- pitates with nitrate of silver.	Crystals tabular, cleavage imperfect.	Matlockite.
			Difficultly soluble in nitric acid.	B. B. with soda easily reduced with the for- mation of a sulphide.	Anglesite.
			Dissolves in hydrochloric acid with separation of PbCl ² to a greenish solution, which, di- luted with water and agitated with tin-foil, assumes a blue color.	fumes escape, and allowed to cool, then breathed upon acquires an ultramarine-	Wulfenite (Molybdate ! of lead.)
			Decomposed by sulphuric acid, leaving a lemon-yellow resi- due. The acid is not colored.	With salt of phosphorus gives in O.F. a colorless glass, which in R.F. becomes blue on cooling.	Stolzite.

T METALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture.	Lustre.	Hard- ness,	Sp. Gr.	Fasibility.	Crystalli- sation.
21 0°.	Brown-yellow.		Resinous.	8.	7.02		17.
² O ⁸ +4aq.	Brown-yellow.	Brittle.	Resinous.	4.	4.7		Amorph
$s^2O^8 + PbCl^2$.	Yellow-brown.		Resinous.	3.5	7.1	L	ш.
Ca) ${}^{3}As^{2}O^{8} + (Pb, Cl^{3}.$	White.		Adamantine.	3.5-4.	5.45	I.	ш.
² O ⁸ +PbCl ² .	White, brown, green.	Brittle.	Resinous.	8.5-4.	6.5-7.1	1.5	ш.
ı) ∇ ²O [¢] .	Bed.		Greasy.	34.	5.7 ·	Easily.	Massive.
μ.	Hyacinth-red.	Prismatic.	Vitreous.	2.5-3.	6.	1.5	v .
² O ⁰.	Hyacinth-red.	Perfect.	Resinous.	38.5	5.7		IV.
	Red.	· · · · · · · · · · · · · · · · · · ·	Dull.	2.—3.		1.	
	Azure-blue.	Prismatic.	Vitreous.	2.5	5.4	Easily.	v.
+PbCl ² .	White.	3 cleavages.	Adamantine.	3.	6.3	Easily.	п.
'+PbCO³.	Greenish- white to yel- low-gray.	Basal per- fect.	Adamantine.	2.5	6.8—7.	Easily.	₹.
۱.	White.	Conchoidal.	Vitreous to adamantine.	3.5	6.4	Easily.	īv.
+3PbCO ³ .	White, yellow- gray.	Prismatic.	Pearly to resinous.	2.5	6.8	1.5	IV.
+3PbCO ³ .	White, yellow- gray.	Basal.	Resinous.	2.5	6.5	1.5	111.
J1².	Colorless white	Prismatic.	Pearly.	2.5	77.1	Easily.	IV.
1º.	Green to yel- low-white.	Basal, im- perfect.	Pearly.	3.	7.2	Easily.	II.
	White.	Conchoidal.	Adamantine.	8.	6.1-6.8	1.5	I V .
04.	White-red, generally yellow.	Octahedral	Resinous.	8.	6.9	1.5	п.
 r4_	Brown, yellow to red.		Resinous.	8.	7.9	8,	ш.

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MINERALS WITHOUT METALLIC LUSTRE.

B. Fusible from 1-5, and non-volatile.

L Yield a metal or a magnetic mass with soda.

DIVISION 2 (concluded).

Drymon 8 (in part).

II. MINERALS WIT

	ead.	General Characters.	Specific Characters,	Speci
	ed.) itobute of		The color is not changed in O. F. ; the others become yellow or colorless.	Vauquel
	Continui Dive a p	B. B. with borax in R. F. give an emerald-green	The nitric solution gives a precipitate or turbidity with nitrate of silver.	Vanadin
mass.	DIVISION 2.—(Continued.) B. with soda on charcoal gtos a globule of lead.	bead.	Does not give the above reactions. B. B. on charcoal with soda gives a zinc coating.	Decheni riety I chite).
magnetic mass.	DIVIS	ald-green color.	Much like vauquelinite. The nitric solution gives with molybdate of ammonia a yellow precipitate (phosphoric acid).	Laxman (Phosy chrom
8	B. B. w		Same as vanadinite : differs in crystalline form.	Descloiz
ne on	-	gummite, Div. 1, p. 89,		24
acon :	toith lue. ve a	Fuses to a black magnetic slag.	Decrepitates and yields much water in the closed tube.	Chenevi
a metallic globule or	s flame, and tott omes violet blue. 1 charooal give a	The nitric solution gives	In the closed tube gives off water and be-	Bayldon
roods, gree	he blowptpe . mmonia beco em alone on per.		In the matrass gives little water (4 per cent.).	OLIVEN
OUN S	or to t s of an of th	with prismatic crystals.	In the matrass gives more water (7 per cent.).	Clinocla
soda on charcoal,	N 3. i blue col an ences or ; most arsenide	D. D. is submy down!	Soluble in ammonia, with separation of car- bonate of lime mechanically mixed with the mineral.	Tyrolite per fr
112202	VISIO autifu tion of deal of	B. B. in matrass decrepi- tate strongly and give much water.	Soluble in ammonia without residue.	Chalcopi lite.
B. B.	DI gree a be on addi ng arsen ittle glol		The fused assay has an alkaline reaction.	Conicha
T	DIVISION 3. Montenet with hydrochloric actd gree a beautiful blue color to the blowphyse flame, and with nitric actd gree a solution tokich on addition of an excess of ammonia becomes riolet blue, a. B. B. on charcoal evolve a strong arsenical odor : most of them alone on charcoal give a white british globule of arsenide of copper.	Does not decrepitate in ma- trass; assumes a smalt- blue color when gently heated.	Loses 22 per cent. on ignition ; soluble in am- monia with the separation of white flocks.	Liroconi
	hydro e a so rroal		Loses 19 per cent. on ignition.	Euchroit
	t with hydro dd give a so on charcoal		Loses only 5 per cent. on ignition.	Erinite.
	Montened nitric ac	actions.	Loses 13 per cent. on ignition.	Cornwal

METALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture,	Lustre.	Hard- ness.	Sp. Gr.	Fusibility.	Crystallina tion.
aC r³O⁹.	Blackish to olive-green.		Adamantine resinous.	2.5-3.	5.6	Easily.	v.
V ² O ⁸ +PbCl ² .	Brown or yel- lowish.		Resinous.	2.75—3.	6.8	Easily.	Ш.
ζn)∇²O ⁶ .	Yellowish-red or ochre-yel- low.		Dull.	8.5	5.6	Easily.	Stalac.
Ju) ⁶ (P,Cr) ⁴ O ¹⁷ .	Pistachio to olive-green.		Vitreous.	8.	5.77	Easily.	v .
°O ¹ .	Olive-brown to black.			3.5	5.8	Easily.	I V .
رهر. ۲۵۶)۶۸۶°O11+3aq.	Dark-green.			4.5	3.93 (?)	Easily.	Massive.
?b)4A82O9+2aq.	Grass to black- ish-green.			4.5	5.35	Easily.	Massive,
s²O°+aq.	Olive-, leek-, blackish- green.	Sometimes fi- brous.	Adamantine to vitreous.	8.	4.1-4.4	2.	IV.
8°O11+3aq.	Dark bluish- green.	Basal.	Pearly to vitreous.	2.5-3	4.19-4.36	2.	٧.
s²O10+9aq.	Apple-verdi- gris green.	Basal.		1.—2.	8.06	Easily.	IV.
s'O13+12aq.	Emerald-grass green.	Basal.	Pearly.	2.	2.5	Easily.	ш.
Ca) ⁴ (As,P,V) ² O ⁹ +	Pistachio to emerald- green.			4.5	4.12	Easily.	Massive.
₼ l) ² (A8, P) ² O ¹¹ +12	Sky-blue to green.		Vitreous.	2—2.5	2. 9	Easily.	♥.
s ² O ⁹ +7aq.	Leek to emer- ald-green.	Prismatio.	Vitreous,	8.5-4	8.89	2.	IV.
s [•] O ¹⁰ +2aq.	Grass to emer- ald-green.		Dull.	4.5	4.04		Amorph.
s ² O ¹⁰ +3aq.	Emerald to verdigris- green.	Amorphous.		4.5	4.16		Amorph.

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MINERALS WITHOUT METALLIC LUSTRE

B. Fusible from 1-5, and non-volatile.

L Field a metal or a magnetic mass with soils.

DIVISION 8 (concluded),

II. MINERALS WITH

	onia		General Characters,	Specific Characters.	Species.										
	h on addition of an excess of amme ad.	•		Gives much water in closed tube, and forms a gray sublimate.	Atacamite										
			solution yields a pre-	Nearly the same reactions.	Tallingite.										
			cipitate of chloride of silver with nitrate of silver.	Sulphuric acid gives a precipitate of sul- phate of lead.	Percylite.										
		5	autor.	Yields no water in the closed tube.	Nantokite										
0.88.	ich	bead		Compare Atlasite, below.											
magnetic mass.	ution tok	odo		Readily soluble in water; the others are not.	Chalcanth (Blue riol).										
or a	td give a sol		B. B. with soda give a sulphuret which on mois- tening blackens silver.	Insoluble in water. The nitric solution gives a white precipitate with nitrate of baryta. In the open tube gives no odor of sulphur- ous acid.											
lobule	On crust coust grow a integration guorence DIVISION 3.—(Continued.) as color to the forme, and inthe nitric acta becomes violet-blue.		oal yiel		Heated in O. F. burns and emits the odor of sulphurous acid.	Covellite									
ullic g				Resembles Brochantite, and has 16 per cent. of water. Brochantite has but 12.	Langite.										
a metu			color to the flame, and to becomes violet. I odor ; most of them on o	the flame, and to becomes violet most of them on o	Easily and quietly soluble	The concentrated HCl solution gives a white precipitate of subchloride of copper on addition of water.									
oal giv	VISION 3 the flam				the flam becom most of t	the flam becom	the flam becom	the flam becom	the flam becon most of	the flam becon	becon most of	most of	most of	in hydrochloric acid.	The hydrochloric solution gives no precipi- tate with water (sometimes effervesces with acids on account of impurities).
char	D1 olor to				Gives much water in closed tube. Color green.	Malachit									
da on	bine c	senical	Dissolve in nitric acid with		Azurite.										
with soda	thing	B, emit no arsenical	effervescence giving off carbonic acid.	Gives with soda a zinc-coating on charcoal.	Aurichalci										
B. wi	a beau			The nitric solution gives with nitrate of silver a precipitate of chloride of silver.	Atlasite.										
В.	B. B	m	Easily and quietly soluble in nitric acid; solutions	Loses 7 per cent. of water on ignition.	Libethenit										
ľ	I Mointened with hydrochioric acts five B	0	give with molybdate of ammonia a yellow preci-		PSEUDOM ACHITE										
			pitate of phospho-molyb- date of ammonia.	Loses 10 per cent. of water on ignition.	Tagilite.										
		d toth hydrochio	d totth hydrochlos	The nitric solution has a yellowish - green color, and gives, with an excess of ammonia, a bluish- green precipitate, and a blue solution.	The solution gives when warmed with mo-										
			the solution obtained ad	a and the mass fused, then boiled with water, cidified with HCl and then boiled down, the reen, and when diluted with water, sky-blue.	Volborthi										

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IETALLIC LUSTRE.

Composition.	Color.	Cleavage or	Lustre.	Hard- ness,	Sp. Gr.	Fusibility.	Crystallina tion.
÷3H²CuO².	Leek-, black-, olive-,emer- ald-green.	Prismatic.	Vitreous.	8.5	4.25	Fusible.	IV.
$+4\mathrm{H}^{2}\mathrm{CuO}^{2}+4\mathrm{aq}.$	Blue to green.			8.	8.5 (?)	Fusible.	Massive.
$\overline{\mathbf{u}}(\mathbf{Cl}^2,\mathbf{O})+\mathbf{aq}.$	Sky-blue.			2.5	-		L
	White.			<u> </u>			
	Sky-blue.			2.5	2.21	Easily.	VI.
) ¹ +8aq.	Emerald to blackish- green.	Prismatic.		8.5-4	3.4	Easily.	IV.
	Indigo-blue- black.	Basal.		1.5-2.	4.6		111.
)•+4aq.	Greenish-blue.			2.5-3	8.5		IV.
	Cochineal-red.	Octahedral.	Earthy, adamantine.	8.5-4.	6.	Easily.	I.
	Black to brownish- black.		Metallic to earthy.	8.	5.95	Difficult- ly.	IV.
) ⁴ +H ² O.	Grass to emer- ald-green.	Fibrous.	Silky to earthy	3.5-4.	3.8	2.	v .
$0^7 + H^2O.$	Blue,		Vitreous.	4.	8.7	2. ·	٧.
1) ³ CO ⁵ +2aq.	Bluish-green,		Pearly.	2.		Difficult- ly.	Acicular.
O ⁴ +CuCl ⁹ +10aq	Celandine to emerald-green		Vitreous-silky	84.	8.85		
0°+H°O.	Dark olive- green.		Resinous.	4.	8.7	2.	I V .
0 ¹¹ +3 aq.	Dark-green.		Vitreous.	4.5-5.	4.2	2,	I V .
0°+3aq.	Verdigris to emerald-green		Vitreous.	8.—4.	4.07	· [•	٧.
?°0''+8aq.	Grass-leek- apple to emer- ald-green.	Micaceous.	Pearly.	2.—2.5	8.5	2.5	п.
a)⁴∇²O°+H²O.	Olive-green to lemon-yellow.	Basal.	Pearly.	8.—8.5	8.5	Easily.	ш.

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MINERALS WITHOUT METALLIC LUSTRE.

B. Fusible from 1-5, and non-volatile.

L Yield a metal or a magnetic mass with sode.

DIVISION 4.

Division 5 (in part).

II. MINERALS WIT

		ead.	General Characters.	Specific Characters,	Spec		
	DIVISION 4.	blue color to a boraz bead.	B. B. in matrass yields much water and becomes smalt-blue.	In HCl soluble to a rose-red solution.	ERYTE (cobalt-		
	DIVI	ue color	Fuses with difficulty, col- ors the flame green.	Soluble in hydrochloric acid, with evolution of chlorine.	Heterog		
	8	ā	The HCl and nitric solu- tions have a green color,	Ammonia gives a green precipitate, which is dissolved in an excess to a sapphire-blue solution.	Annab (always tains a coba		
a magnetic mass.	visions.	evolve a strong odor.	Fuse easily B. B. to mag- netic beads. The color of the pulverized miner-	Amorphous.	Pitticit		
gneti	the di	volve ;	als are quickly changed to reddish-brown by a	Crystallization isometric.	Pharma rite.		
a ma	but do not give the reactions of the preceding divisions.	nical e	solution of caustic pot- ash.	Crystallization orthorhombic.	SCOROL		
le or		a) During fusion e arsenical	Fibrous, with silky lustre.	Color brownish-yellow-9 p. c. water.	Arseni		
globu	tions	a) Du	Mostly soluble in water.	With excess of ammonia gives a blue solu- tion. Sometimes contains arsenic.	Moreno		
a metallic globule	the the react	ng.	Gives much water (13 p.c.) in the closed tube, and col- ors the borax-bead blue.	Soluble in strong HCl with evolution of chlorine. Soluble in phosphoric acid to a violet fluid.	Rabdio		
dive a s	not git	HCI without leaving a perceptible residue, and without gelatinizing	Gives antimonial fumes on charcoal.	Gives water in the closed tube.	Stibiof		
oal g	op m	gelati		Gives little or no water in the closed tube.	Pettkoi		
with soda on charcoal	5. 10188, b	rithout		Perfectly soluble in water.	MELA ITE (CO		
a on	DIVISION 5. gray magnetic mass.	and w	B. B. swell up and in the	B. B. swell up and in the	B. B. swell up and in the	Soluble in water, leaving a yellow residue.	Botryo
11 800	DIV U mag	esidue,	R. F. fuse perfectly to a magnetic slag. The so- lutions give with chloride	(Roeme		
B. wat	or gra	tible r		Same reactions as Botryogen. Their pow-	Coquin		
B. I	black o	percep	ryta, and with ammonia a greenish precipitate,	red by solution of potassa.	Jarosit		
T	diec a	ing a	which in the air changes to brownish-red; all ex-		Fibrofe		
3	R. F.	ut leav	cept Pettkoite, give much water in the closed tube.	[COPIAL		
	in the	witho		Insoluble in water; powders yellow.	Raimon		
	rcoal	n HCI			Carpho		
	on che	Soluble in		Characterized by its color and octahedral crystallization.	Voltait		
	B. fused on charcoal in the R. F. give a black or	b) So	Soluble in heated HCl with effervescence.	Difficultly fusible ; becomes by heating black and magnetic.	Siderit (spathi		
	B. B.	-	enervescence,	Compare Mesitite, Div. 4, page 92. Compare Sphalerite.	S		

IETALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture.	Lustre,	Hard- ness.	Sp. Gr.	Fusibility.	Crystallina tion.
² O ⁸ + 8aq.	Crimson, peach-red.	Prismatic.	Pearly.	1.52.5	2.94	2.	v.
+2Co ² O ³) + 6aq.	Black—red- brown.			3.		Difficult- ly.	Amorph.
O ⁸ +8aq.	Apple-green.		Earthy.	Soft.		Easily.	v .
As ² O ⁵ SO ³ ,H ² O.	Yellow to red- dish-brown.			2.—3.	2.2-2.5		
⁶ O ²⁷ + 15aq.	Green-red- brown, yellow.	Cubic.		2.5	3.	Easily.	L.
0 ⁸ + 4aq.	Leek-green to brown.			8.5-4.	3.2	Easily.	IV.
$Fe)As^2O^8 + H^6FeO^6$	Brownish-yel- low.		Silk y .	1.—2.	8.8	Easily.	
+7Aq.	Apple-green to bluish-green.		Vitreous.	2.	2.		
[n,Co)(Fe, M n)O⁴.	Black.			Soft.	2.8	3.	
,Sh ² O ⁵ ,H ² O.	Yellow.		Resinous.	4.	8.52		I V .
² e)S ³ O ¹⁹ .	Black.			2.5		Fuses.	
+ 7aq.	Green.			2.	1.8	-	v .
[g)FeS4016+12aq.	Ochre-yellow to red.			2.5	2.04		٧.
n)FeS4O16+12aq.	Yellowish- brown.			2.75	2.17	-	v.
)19 + 9aq.	White to yel- low.			2.5	2.		ш.
S4O29 + 6aq.	Ochre-yellow.		-	8.	3.2		пі.
0°+10aq.	White to pale yellow.			1.5	1.84		Fib.
O ⁹¹ +18aq.	Sulphur - yel- low.		Pearly.	1.5	2.14		ш.
O ¹⁵ +7aq.	Honey- to ochre-yellow.		Pearly.	8.	3.19		ш.
0 ⁹⁷ + 13aq.	Straw-yellow.		Resinous.	4.	2.5		Mass,
⁸ O ¹³ + 20aq.	Black dark- green.		Resinous.				L.
, ³ .	Ash-gray to brownish-red.	Rhombohe- dral.	Pearly-vitre-	4.	8.6-8.9	4.5	Ш.

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MINERALS WITHOUT METALLIQ LUSTRE.

B. Fusible from 1-5, and non-volatile.

L Yield a metal or a magnetic mass with sode.

Drymon 5 (continued).

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II. MINERALS W

				General Characters.	Specific Characters.	Species.		
	(latelone.	gelatinizing	monia a yellow precipi-	Gives water in the closed tube.	Hureaulite.		
		preceding o	nd without		Gives the reaction for fluorine when fused	TRIPLITE.		
atue.		DIVISION E (Continued.) or gray magnetic mass but do not give the reactions of the preceding divisions.	residue a	glass (manganese).	Distinguished from triplite by its color.	Sarcopside.		
Fusible from 1b, and not volatile, or only partially volatile.	with soda on charcoal give a metallic globule or a magnetic mass.		b) (Continued.) Soluble in HCl without leaving a perceptible residue and without gelatiniatag.	Gives the above phosphoric acid reaction with mo- lybdate of ammonia. The blowpipe flame is colored purple-red in streaks (lithia).	With borax gives the manganese reaction, but not so plainly as the minerals of the	TRIPHYLITE		
r only	or a me		out leavin	Gives the above phosphoric The solution with chlor sulphate of baryta.	acid reaction with molybdate of ammonia. ide of barium yields a heavy precipitate of	Diadochite.		
intie, o	globule	DIVISION 5.— (Continued.) - gray magnetic mass bu	HOI with	Give the above phosphoric acid reaction with mo- lybdate of ammonia.	Loses 28 per cent. of water on ignition.	Vivianite.		
VOIa	tallic	-(Co	le in J	Moistened with sulphuric acid the flame is colored	Loses 10 per cent. on ignition.	DUFRENITE.		
DOL	a me	v mag	Solut	pale-green. Easily fusi- ble. The borax - glass	Loses 19 per cent. on ignition.	Borickite,		
and	give	DIVIB T gra	nued.)	shows only the reactions for iron.	Loses 33 per cent. on ignition.	Cacoxenite.		
	sharcoal	F. gtve a black o	b) (Conti		Tompare Beraunite.	Beraunite.		
8	no	gtee		Difficultly fusible to mag-	Streak red.	Hematite.		
210	soda	B. Juset on charcoal in the R. F.	in HCl, forming a jelly, or with the separation of allica.	r with	r with	netic globules.	Streak yellow.	LIMONITE.
23 I.S.						r with	In a matrass yields water, and with HCl forms a perfect jelly.	Fuses with slight puffing to a black glass.
1	L. B. B.	charcoal			Radiated, sometimes foliated.	STILPNOME- LANE (Chai codite).		
		uo 1		Cl, forming separation o	In the matrass yield wa-	Micaceous,	Voigtite.	
		L. Juse			Cl, for	Cl, for separat	ter, and are decomposed by HCl without gelatin-	Massive.
		B. J	e in H the	izing.	Massive.	Euralite.		
			e) Soluble		Sometimes gelatinizes, sometimes does not.	Palagonite.		

JUT METALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture.	Lustre,	Hard- ness,	Sp. Gr.	Fonblity.	Orystalliss tion.
'e,H')3P'O8+4aq.	Orange to red- dish-yellow.		Vitreous.	5.	8. 2	Easily.	v .
n) ³ P ² O ⁸ +(Fe,Mn)	Brownish- black.	Cleavable in three di- rections.	Resinous	5.	3.6	1.5	IV.
Fe)*P*O* + H*Fe	Flesh-red to lavender- blue.		Silky.	4.	8.7	1.5	V. (?)
n,Li²)²P²O ^{\$} .	Greenish-gray, bluish, etc.	Perfect.	Resinous.	5.	8.54	1.5	I V .
SO ³ , P ² O ⁵ , H ² O.	Red, yellow, brown.	Brittle.	Resinous.	8.	2.03	Easily.	Amor- phous.
) ⁸ +8aq.	Different shades of blue.	Perfect.	Pearly-vitre- ous.	1.5-2.	2.6	1.5	٧.
) ¹¹ +3aq.	Dark leek- green.	Radiated.	Silky.	3.5-4.	8.3	Easily.	I V .
³) ⁵ P ⁴ O ²⁵ +15aq.	Reddish- brown.		Waxy.	3.5	2.7	Easily.	Mass.
) ¹¹ +12aq.	Brownish yel- low.	Fibrous, ra- diated.	Silky.	3-4.	3.38		
⁸ +aq.	Hyacinth-red, reddish- brown.	Foliated.	Metallic, pearly.	2.	2.87	Easily.	
	Reddish black.	Compact, earthy.	Dull.	66.5	4.5	5.	111.
+3aq.	Brown.	Compact, fibrona,	Dull.	δ.	3.6 4	5.	
$fg)^{2}SiO^{4} + Fe^{2}Si$ -6aq.	Raven-black.	Basal.	Vitreous.	3.5	3.35	Easily.	III.
;) ² (Fe, Al)Si ⁵ O ¹⁵ +	Bronze-yellow to greenish- gray.	Radiated, compact.	Pearly to sub- metallic.	3.	2.76	Easily.	
Mg, Si, aq.	Leek-green, yellow.	Micaceous.	Pearly.	2.5	2.91	Easily.	
, F eŜi,aq.	Leek-green to black.		Greasy.	2.5		Easily.	
lfeŜi,aq.	Dark-green to black.			2.5	2.62	Easily.	
Mg,Ċa,Ŝi,aq.	Yellow-red, black.		Vitreous, greasy.	45.	1.8-2.7	Es-ily.	

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MINERALS WITHOUT METALLIC LUSTRE.

B. Fusible from 1-5, and non-volatile.

L. Field a metal or a magnetic mass with sode.

Drymon 5 (cominned).

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11. MINERALS WIT

			General Characters.	Specific Characters.	Speci				
				Intumesces slightly; decrepitates slightly; fuses quietly to a black magnetic bead.	ILVAITE				
88.	give the reactions of the preceding divisions.	separation of silica.	Give little or no water in the closed tube; with HCl they gelatinize. Not cleavable.		Allani				
tic ma	na of t	the		Magnetic.	FATALI				
a magnetic mass.	the reactio	lly, or with	or	Crystalline and cleavable;	Decomposed by phosphoric acid the jelly immediately becomes violet when treated with nitric acid.	HORTON LITE.			
04		forming a jelly,	gelatinize perfectly.	Same reaction.	Knebeli				
globu	(') # qo n			Gives with soda a sublimate of oxide of zinc.	Roepper				
netallic	onkinue	e in HCI,	Decomposed with separa-	Mixed with salt of phosphorus and oxide of copper tinges the flame green (chlorine).	Pyrosm				
1 20 00.	r b(C				Soluble		tion of silica without gelatinizing. $F=2$.	The HCl solution, boiled with tin, is colored violet (titanic acid).	Astroph
charcoul give a metallic globule	DIVISION 5 (Continued.) or gray magnetic mass, but do not	(Continued.)	Decomposed easily by HCl, leaving a residue of sil- ica in the form of scaly flakes.		LEPIDO LANE				
ano	a black	()	In some varieties forms an imperfect jelly with HCl.	Not cleavable, easily fusible.	ALLOCH (iron garne				
with soda	Bang. R. Bue		magnetic. Decomposed	Amorphous.	Gillingi				
B. B.	the		by HCl without forming a jelly. Give water in matrass.	Fibrous.	Xylotile				
н	2001					Compare Limonite, Div. 4, p. 92.	1.		
	a char	ydro-	Tinges the flame purple- red (lithia).	Very perfectly cleavable in one direction (micaceous).	Lepido				
	fused on charcoal in	upon by hydro-	n by h	Imparts a violet color to the borax bead.	Some specimens contain enough iron to be- come magnetic.	Rhodo			
	B.B.J	acted ric ac	Decomposed by aqua-regia with separation of a	give the bluish-green manganese reaction.	WOLFR				
		vitightly	yellow powder (tungstic acid).	Same reactions.	Megaba				
		y stig	wont).	Same reactions, contains no iron.	Hübner				
		d) Only	Fuses quietly at 3. Gela- tinizes after fusion.	Not easily cleavable.	Alman Garr				

ETALLIC LUSTRE.

Jomposition.	Color.	Cleavage or Fracture.	Lustre,	Hard- ness,	Sp. Gr.	Fusibility.	Crystalli zation.
^r e ⁴ FeSi ⁴ O ¹⁸ .	Gray to iron- black.			5.5—6.	4.	2.5	IV.
., Di, Fe, Ca) ² (Al i ³ O ¹³ .	Pitch-brown to black,		Pitchy to res- inous.	5.56.	8.—4.2	5.5	I V .
۹.	Dark-green brown to black.	Two cleav- ages at 90°.	Resinous.	6.5	4.	Easily.	
;)²SiO4.	Yellow to dark-yellow green.	Three cleavages.	Resinous.	6.5	8.91	4	IV.
ı) *SiO 4.	Gray, red. brown to black.			6.5	4.12	Easily.	IV.
ı,Zn,Mg) ² SiO ⁴ .	Dark-green to black.	Rectangu- lar.	Vitreous.	6.	4.	Diff.	IV.
n)Cl ³ + 7(Fe,Mn)) + 5aq.	Brown to blackish- green.	Basal.	Pearly.	4.5	8,16	2.5	ш.
) ⁶ (Fe, Mn) ¹⁵ (Fe, Si,Ti) ¹⁶ O ⁵⁶ .	Bronze-yel- low.	Micaceous.	Pearly.	3	3.32	Easily.	IV.
.	Dark-green to black.	Micaceous.	Vitreous.	8.	8.	Easily.	111.9
3i ³ O ¹³ .	Green-yellow to black.		Greasy.	7.	3.7—4.	8.	L.
Mg,Ċa,Ŝi, a q.	Brownish- black.		Dull.	3.	8.04	•	Amor- phous.
, Ŝi, aq.	Wood-brown.	Fibrous.	Silky.		2.4		
Li) ⁴ Al4Si ¹² O ³⁹ .	Rose-red, gray-white.	Micaceous.	Vitreous.	2.5—3.	2.8-3.	2.—2.5	I V .
².	Bose-red, brownish- red.	Perfect.	Vitreous.	5.5—6.5	3.61	2.5	VI.
a)WO4.	Black.	Prismatic.	Sub-metallic.	5.—5.5	7.1-7.5.	2.5-8	♥.
e)WO '.	Brown.	Prismatic.	Sub-metallic.	8.5-4.	6.4-6.9	Easily.	v .
μ.	Brown-red.	Prismatic.	Adamantine.	4.5	7.1	Easily.	v .
3i ² O ¹⁹ .	Red or brown- ish-red.		Vitreous.	77.5.	3.7-4.	8.	L

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MINERALS WITHOUT METALLIC LUSTRE

B. Fusible from 1-5, and non-volatile.

I. Field a metal or a magnetic mass with soda.

DIVISION 5 (concluded).

Devanour 4.

II. MINERALS WITH

	ugnetic na.	vold.	General Characters.	Specific Characters,	Species.	
magnetic mass.) ack or gray mo eceding divisio	upon by hydrochloric a	Fuse quietly to a black shining glass.	Fused with soda, and then dissolved in HOI and treated with ammonia to separate iron, the filtrate gives with oxalate of ammonia a heavy precipitate (lime).	Pahinatonina	
metic	5(Continued.) R. F. give a bla actions of the pri-			Gives no lime when treated as above.	Acmite.	
ma	P. Co	octed		Compare Augite, Div. 6, p. 88.		
r a	No R.	htly :	Easily fusible (1.7-2) with strong intumescence and	Gives water in the closed tube.	Crocidolite.	
bule o	DIVIBIO OGI In I	B. B. fused on clu mass, but do no d. (Continued.)	escape of gas bubbles to a black glass.	Yields no water in a matrass.	ARFVEDSON- ITE.	
metallic globule or	I on churo do not g		Fuses at 3 without swel- ling. Gives water in matrass.		Glauconite (Green earth	
give a	B. B. fused mass, but		d. (Contin	d. (Contin	Compare Amphibole Div. 6, p. 88, Tourmalins, Div. 6, p. 87. Compare Le- pidomelane, Subdivision a, p. 78.	
on churcoal			Easily soluble in HCl, yielding a colorless solu- tion, which becomes blue on agitation with tin- foil.	sorbed. In R. F. with salt of phosphorus	Molybdite.	
B. with soda	oregoin		1.000	Gelatinizes perfectly with HCl.	Eulytite,	
3. with	DIVISION 6.	vistons.	Fused with sulphur and iodide of potassium on charcoal give a fine red		Bismutite.	
I.B.I	DIVISION 6. Not belonging to the foregoing	di	sublimate on the coal (bismuth). Compare Walpurgite, p. 82.	With salt of phosphorus gives a grean head	Pucherite.	
	×		Compare Samar- skite, Div. 6, p. 69; Allan- ite and Lepidomelane, Div. 5, p. 78.			

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IETALLIC LUSTRE

omposition,	Color.	Cleavage or Fracture.	Lustre,	Hard- ness.	Sp. Gr.	Fusibility.	Crystalliza tion.
., M n)SiO ³ + F e	Dark green- black.		Splendent.	5—6.	8.86	2.5	VI.
F e)Si ³ O ⁹ .	Red-brown to blackish- green.	Cleavable at angle of 93°.	Vitreous.	6.	8.4	2.	v.
Mg,Ŝi,aq.	Green to lav- ender-blue.	Fibrous.	Silky.	4.	3.2	Easily.	
e,Ca)SiO ^s + Fe		Perfectatan angle 123°.	Vitreous.	6	3.4	2.	v.
È,Äl,Ši,aq.	Deep-olive to sea-green.	Scaly.	Dull.	Soft.	1.—2.	2.2-2.4	
	Sulphur- orange, yel- low.		Silky, earthy.	1.—2.	4.5	1.	τ ν .
	Dark hair- brown to yellow.		Resinous.	4.5	6.1	Easily.	 I.
¹⁸ +9H ² O.	White to yel- low.		Dull.	44.5	6.8—7.6	Easily.	Amorph.
	Reddish- brown.	Basal.	Vitreons adamantine.	4.	5.91	Easily.	IV.
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MINERALS WITHOUT METALLIC LUSTRE.

B. Fusible from 1-5, and non-volatile.

L. Vield no metal or magnetic mass with sods

Nummer 1 (in part).

II. MINERALS WIT]

	-brown		General Characters,	Specific Characters,	Specie					
	molstened turmeric paper to red brow		B. B. on charcoal defla-	Fused on platinum wire colors the flame violet. In the solution bichloride of pla- tinum produces a yellow crystalline pre- cipitate.	Nitre.					
	ed turmes		grate strongly.	Fused on platinum wire colors the flame strongly yellow. Bichloride of platinum produces no precipitate.	Soda Ni					
в.	olsten		In a matrass yield much	Rapidly effloresces on exposure to the air and changes to thermonatrite.	Natron.					
111.03	n of m		water; the aqueous so- lutions react alkaline,	Effloresces.	Thermon trite.					
magnetic mass.	the color	the color		and effervesce on addi- tion of an acid.	Does not alter on exposure.	TRONA.				
globule or m	and change		10	The solution gives a white precipitate with soda. Ignited and treated with cobalt solution yields a flesh-red mass (50 per cent. water).						
tallic gl	etallic g 1. 1. reaction,	in water.		With soda yields a white precipitate. Ignit- ed and treated with cobalt solution yields a blue mass.	KALIN (potash a					
give no metallio Dryston 1. an alkaline reactio	Easily and completely soluble in water.	The aqueous solution does	In the concentrated solution bichloride of platinum yields a yellow precipitate. Not affected by the above reagents; yields							
oal g	have a	have a	have a	have a	omplet	omplet	omplet		water in the closed tube. Not affected by the above reagents; yields	(glauber Thenard
charcoal	reepa	ando	Chloride of barium gives an abundant white pre- cipitate of sulphate of	no water in the closed tube. Like epsomite—14 per cent. water.	Loeweite					
a on	a the fe	Easily a	Easily	baryta, which is insolu- ble in acids.	Like epsomite—13 per cent. water.	Kieserite				
with soda	oal or in	(u)		Like epsomite—21.5 per cent. water.	Bloedite.					
B. B. w	n charo			Like epsomite but does not effloresce in air.	Simonyit					
П. 1	ating o			Like epsomite—loses 26.8 water when heated to 133° C.	Picromer					
	ontinued he		Yield no precipitates in the aqueous solutions with chloride of barium or alkalies; with nitrate of	Yields a heavy precipitate with bichloride of platinum.	Sylvite.					
	Drvision 1. B. after fusion and construed heating on charcoal or in the forceps have an alkaline reaction, and change the color of a) Easily and completely soluble in wrter.		silver yield a heavy pre-	Yields no precipitate with bichloride of pla-	Hali (common					
	B. B. after		Moistened with strong sul- phuric acid gives a green flame (boric acid).	Reaction alkaline; does not effervesce with acids; bubbles, swells up, and fuses to a clear bead B. B.	Borax.					

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ETALLIC LUSTRE

Domposition.	Color.	Cleavage or Fracture.	Lastre.	Hard- bons.	Bp. Gr.	Fasibility.	Cryvia satis
	White.		Vitreous.	8.	1.98	Easily.	IV.
i.	White.		Vitreous.	1.5-2	8.8	Easily.	111.
*+10aq.	Gray-white.		Earthy.	1.5	1.4	Easily.	v .
⁸ + aq.	Gray-white.			1.5	1.5-1.6	Easily.	IV.
)8+8aq.	Gray-white.			2.5-8.	2.11	Easily.	v .
+7aq.	Colorless- white.		Vitreous.	2.25	1.7	Easily.	IV.
40 ¹⁶ +24aq.	White.		Vitreous.	2.25	1.75	Easily.	I.
	White.		Vitreous.	8.	1.78	Easily.	IV.
4+10aq.	White.		Vitreous.	1.5-2.	1.48	Easily.	v .
4.	White.		Vitreous.	2.—3.	2.55	Easily.	IV.
gS ² O ⁸ + 5aq.	Yellow-white, red.		Vitreous.	2.5	2.87	Easily.	II.
+ aq.	White.		Dull.	2.5	2.51	Easily.	IV.
S'0 ² + 4aq.	White, orange- red.			_		Easily.	v .
$S^2O^3 + 4aq.$	Colorless to blue-green, yellow.			2.5	2.24	Easily.	v.
5°O ⁸ +6aq.	White.		Silky.	2.5		Easily.	v
	Colorless to white.	Cubic.	Vitreous.	2.	1.9—2.	Easily.	I .
	Colorless, white, red, purple.	Cubic.	Vitreous,	2.5	9.15	Easily.	L.
0'+10 aq.	Gray-white.		Vitreous.	2.5	1.72	Eastly.	v .

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MINERALS WITHOUT METALLIC LOS'TRE.

B. Fusible from 1-5, and non-voiatile.

IL Yield no metal or magnetic mass with sode.

DEVERSOR 1 (conclusied).

II. MINERALS WITHO

			General Characters,	Specific Characters.	Speci															
	alkaline reaction and change the color of motstened turmeric paper		Fusibility=1. Alone colors the flame yellow; mois- tened with strong sul- phuric acid colors the flame green (boric acid).	soluble in hot water; the solution is alka-	ULEXIT															
	ened			Gives much water in the closed tube.	Gay-luss															
	of moist		Soluble in dilute hydro- chloric acid with effer-	The dilute solution gives a heavy precipitate with sulphuric acid; fused in the forceps, colors the flame green.	Wither															
with sode on charcoal give no metallic globule or magnetic mass.	uge the color		vescence.	The HCl solution gives a precipitate with ammonia (phosphate of lime). The nitric solution, warmed with molybdate of am- monia, gives a yellow precipitate.	Staffalit															
magn	nd char		Quietly soluble in much	In closed tube yields much water.	Gypsun															
obule or	eaction a	water.	hydrochloric acid; the solution gives with chlo- ride of barium an abun-	Yields little water in the closed tube. In its solution bichloride of platinum gives a yel- low precipitate. Partially soluble in water.	POLYHA															
tallic gl	ikaline r	an on	charcoal or in the forcept, have an alkaline i to real-brown. b) Insoluble or difficulty soluble in w	in the forceps, have an alkaline to red-brown. Insoluble or difficultly soluble in w	in the forceps, have an alkaline i to red-brown. insoluble or difficulty soluble in w	tralized by ammonia	Yields no water, and is not precipitated by bichloride of platinum. Partially soluble in water.													
e no me	stor 1 zre an a -brown. althy solu					n the forceps, have an a to red-brown. nsoluble or difficultly solu	n the forceps, have an al to red-brown. nsoluble or difficultly solul	n the forceps, have an al to red-brown. nsoluble or difficultly solul	seps, have an al to red-brown. or difficultly solu	ave an a -brown. ruitly solu	rve an a -brown. althy solu	rve an a- brown. althy solu	ave an a -brown. cultly solv	ave an a t-brown. cultly solt	debrounn.	de an c d-brown. cultly soli	d-brown. leultly sol	gives with oxalate of ammonia a precipitate of oxalate of lime.		Anhydr
al giv	DIVISION eps, have (to red-bros										Compare <i>Celestite</i> , below, which in fine powder is slightly acted on by acids.									
charco	n the forc								Very little acted upon by HCl. B. B. with soda	Fused in the forceps colors the flame yellow- ish-green.	Barite.									
la on	t or t			give a sulphur reaction.	Fused in the forceps colors the flame red.	Celestit														
with soc	charcoa			9	Heated on charcoal evolves an arsenical odor.	Yields water in a matrass.	Pharmac													
B.	no Dui			Easily fusible in the flame of a candle. $(F = 1.)$	Cryolite															
IL B.	d heat		1.	In the closed tube decrepitates and generally phosphoresces.	Fluor (fluor s															
-	linue		When fused with bisul- phate of potassa in a	Same as cryolite (occurs in granular masses).	Chiolite.															
	noo but		matures might manus of	The same, but in closed tube yields water, which has a strongly acid reaction.	PACHNO (Thomse															
	NON C		corroue the glass,	Yields no water in the closed tube.	Arksutite															
	r Ju			Yields no water in the closed tube.	Chodneff															
	afte			Yields water in the closed tube.	Gearksut															
	B. B.		Effervesces with concen- trated hydrochloric acid; the solution when eva- porated gelatinizes.	B. B. immediately becomes white and opa- que; fuses at 2.5 with intumescence to a white blistered glass, which placed on tur- meric paper gives an alkaline reaction.	CANCERD (near ne lite)															

CALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture.	Lustre.	Hard- ness.	Sp. Gr.	Fusibility.	Crystalliza tion.
B509+5aq.	White.	Fibrous.	Silky.	1.	1.65	1.	
) ³ +CaCO ³ +5aq.	White.		Vitreous, pearly.	2.—3.	1.99	Easily.	v. •
3.	White-gray.		Vitreous.	8.5	4.3	2.	IV.
0 ⁸ +CaCO ⁸ .	Leek-green to green-yel- low.			4.	8.18		Stalact.
+2aq.	Colorless, gray-white.	In 3 direc- tions.	Silky, vitreous.	2.	2.3	2.5—3.	v .
K ² S ⁴ O ¹⁶ +2aq.	Yellow to brick-red.		Vitreous.	2.5	2.77	1.5	IV.(?)
S²O ⁸ .	Yellow to gray.		Vitreous.	2.5	2.7	1.5	v .
	Colorless, white-blue, red.	Perfect in 3 directions.	Vitreous.	8.5	2.9	2.5-3.	IV.
·	All colors, white - yellow, blue.	Basal, per- fect.	Vitreous.	 2.5—3.5	4.5	8.	IV.
	Colorless, white, blue.	Basal, per- fect.	Vitreous.	3.—3.5	3.9	3.	I V .
\sO⁴+5 aq .	White-gray.		Vitreous.	2.—2.5	2.7	Easily.	v.
F19.	White to black.	Basal per- fect.	Vitreous.	2.5	3.	1.	17.9
	All colors.	Octahedral.	Vitreous.	4.	8.18	3.	I.
F'.	Snow-white.			4.	2.72	I.	ц
' A lF ¹² +2 a q.	Colorless- white.		Vitreous.	2.5-4.	2.75	Easily.	v .
AlF ¹⁰ .	White.		Vitreous.	2.5	8.1	Easily.	Mass.
E10	White.			4.	3.	Easily.	<u>II.</u>
? ¹⁰ +4aq.	White.		Earthy.	2.			
Si ² O ⁸ som e carbonic).	White, pink, gray-yellow.	Hexagonal.	Vitreous.	5.—6.	2.5	2.5	II I.

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MINERALS WITHOUT METALLIC LUSTRE

B. Fusible from 1-5, and non-volatile.

II. Yield no metal or magnetic mass with sode.

Devenuos 2 (in part).

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MINERALS WITHOUT METALLIC LOS'TRE.

B. Fusible from 1-5, and non-voiatile.

U. Yield no metal or magnetic mass with sods.

DEVENION 1 (conclusied).

II. MINERALS WITHO

			General Characters,	Specific Characters.	Speci													
	turmeric paper		Fusibility =1. Alone colors the flame yellow; mois- tened with strong sul- phuric acid colors the flame green (boric acid).	soluble in hot water; the solution is alka-	TIEXIT													
	ened			Gives much water in the closed tube.	Gay-luss													
	of motes		Soluble in dilute hydro- chloric acid with effer-	The dilute solution gives a heavy precipitate with sulphuric acid; fused in the forceps, colors the flame green.	Witheri													
on charcoal give no metallic globule or magnetic mass.	uge the color		vescence.	The HCl solution gives a precipitate with ammonia (phosphate of lime). The nitric solution, warmed with molybdate of am- monia, gives a yellow precipitate.	Staffelit													
magn	nd cha		Quietly soluble in much	In closed tube yields much water.	Gypsum													
obule or	eaction a	alkaline reaction and change the color of motstened turmeric paper while in writer.	alkaltne reaction o able in water.	hydrochloric acid; the solution gives with chlo- ride of barium an abun-	Yields little water in the closed tube. In its solution bichloride of platinum gives a yel- low precipitate. Partially soluble in water.	POLYHA												
stallic gl	ikaline r			alkaline ; uble in w	alkaline Inble in w	luble in w	tralized by ammonia	Yields no water, and is not precipitated by bichloride of platinum. Partially soluble in water.	GLAUBE									
ne no me	Toe an a	or difficultly soluble	gives with oxalate of ammonia a precipitate of oxalate of lime.	Yields no water; does not precipitate by bichloride of platinum; insoluble in wa- ter.	Anhydr													
al giv	DIVISION eps, have to	B. after fusion and continued heating on charcoal or in the forceps, have an is real-brough. b) Insoluble or difficulty so	or in the for b) Insoluble	seps, h	to red r diffic	to red r diffic	eps, ho to red r diffic	eps, ho to red. r diffio	eps, ho to red. r diffic	eps, ho to red r diffic	eps, ho to red r diffic	to red	seps, ho to red or diffic	eps, ho to red r diffic	or diffic		Compare Celestite, below, which in fine powder is slightly acted on by acids.	1
charco	a the forc			Very little acted upon by HCl. B. B. with soda	Fused in the forceps colors the flame yellow- ish-green.	Barite.												
ta on	d or it			5 6	6) OL	give a sulphur reaction.	Fused in the forceps colors the flame red.	Celestite										
with soda	charcoa			Heated on charcoal evolves an arsenical odor.	Yields water in a matrass.	Pharmac												
B. B. u	no Oui			Easily fusible in the flame of a candle. $(F = 1.)$	Cryolite													
II. J	d hea			In the closed tube decrepitates and generally phosphoresces.	Fluor (fluor s)													
7	sinue		When fused with bisul- phate of potassa in a	Same as cryolite (occurs in granular masses).	Chiolite.													
	noo pur		matrass, yield vapors of hydrofluoric acid, which corrode the glass.	The same, but in closed tube yields water, which has a strongly acid reaction.	PACHNO (Thomse													
	nois		corroue the glass.	Yields no water in the closed tube.	Arksutite													
	er fu			Yields no water in the closed tube.	Chodnef													
	alte .			Yields water in the closed tube.	Gearksut													
4	B. B.		Effervesces with concen- trated hydrochloric acid; the solution when eva- porated gelatinizes.	B. B. immediately becomes white and opa- que; fuses at 2.5 with intumescence to a white blistered glass, which placed on tur- meric paper gives an alkaline reaction.	CANCRI (near ne lite)													

FALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture.	Lustre.	Hard- ness.	Sp. Gr.	Fusibility.	Crystalliz tion.
.B⁵O ⁹ +5aq.	White.	Fibrous.	Silky.	1.	1.65	1.	
	White.		Vitreous, pearly.	2.—3.	1.99	Easily.	v. •
)3.	White-gray.		Vitreous.	8.5	4.8	2.	IV.
'0 ⁸ +CaCO ⁸ .	Leek-green to green-yel- low.			4.	3.18	_	Stalact.
4+2aq.	Colorless, gray-white.	In 3 direc- tions.	Silky, vitreous.	2.	2.3	2.5—3.	v .
3 ^K ² S ⁴ O ¹⁶ +2aq.	Yellow to brick-red.		Vitreous.	2.5	2.77	1.5	IV.(?)
	Yellow to gray.		Vitreous.	2.5	2.7	1.5	v .
4.	Colorless, white-blue, red.	Perfect in 3 directions.	Vitreous.	8.5	2.9	2.5-3.	IV.
٠	All colors, white - yellow, blue.	Basal, per- fect.	Vitreous.	2.5—8.5	4.5	8.	IV.
•	Colorless, white, blue.	Basal, per- fect.	Vitreous.	3.—3.5	3.9	8.	IV.
AsO ⁴ +5 aq .	White-gray.		Vitreous.	22.5	2.7	Easily.	v .
F ¹⁹ .	White to black.	Basal per- fect.	Vitreous.	2.5	3.	1.	17.9
	All colors.	Octahedral.	Vitreous.	4.	8.18	3.	I.
F°.	Snow-white.			4.	2.72	I.	ц
² AlF ¹² +2aq.	Colorless- white.		Vitreous.	2.5-4.	2.75	Easily.	v .
▲IF ¹⁰ .	White.		Vitreous.	2.5	8.1	Easily.	Mass.
F ¹⁰ .	White.			4.	3.	Easily.	<u>.</u>
F ¹⁰ +4aq.	White.		Earthy.	2.			
Si ² O ⁸ some carbonic).	White, pink, gray-yellow.	Hexagonal.	Vitreous.	5.—6.	2.5	2.5	т.

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MINERALS WITHOUT METALLIC LUSTRE

B. Fusible from 1-5, and non-volatile.

II. Yield no metal or magnetic mass with sode.

Devenour ? (in part).

	General Characters.	Specific Characters.	Specie
		Fuses easily; with strong sulphuric acid it gives off hydrofluoric acid, which corrodes glass.	Durangit
		Gives an amethystine bead with salt of phos- phorus (oxide of manganese).	Chondra: ite.
	Give arsenical fumes on charcoal.	Gives a green bead with salt of phosphorus (oxide of uranium; with S + KI gives a red sublimate on charcoal (iodide of bismuth).	
j.		Gives a green bcad with salt of phosphorus, but no reaction for bismuth.	Trögerite
mass. vaporation		Gives on charcoal a coating of oxide of zinc.	Adamite.
II. B. B. with solid on charcool give no metallic globulo or magnetic mass is in hydrochloric acid, some also in water. The solution is not petatinised by staportic	Soluble in water. Colors the borax bead violet when hot (oxide of manganese)	Gives much water (40 p. c.) in the closed tube.	Fauserite
or magnetio gelatinteed by	Soluble in water. Give a sulphur reaction with	8	Tscherm (ammo alum
iobula o is not o	soda on charcoal. Fuse when first heated, and swell up to an infusible		Alunogen
etallio g solution	mass.	After fusion moistened with nitrate of co- balt and again ignited becomes green (oxide of zinc).	Goslarite vitrio
jive no me Divi∎ios 2. ater. The	Treated with caustic pot- ash or soda gives the odor of ammonia.	Gives much water in a matrass.	STRUVIT
oal give Divi n water.		Imparts a violet color to the hot borax bead (oxide of manganese).	Sussexite
aro		Soluble in water.	Sassoli (boric ad
t on charcod some also in	buildesochoo, and coror		Hydrobor
soda acid,		Gives little or no water.	BORACIT
with horte	acid reaction with sul- phuric acid and alcohol.	Like Hydroboracite, but contains only 7 p. c. water.	Szaibelyi
	Compare Boraz, Div. I., p. 80.	Like Hydroboracite. Its nitric solution gives a yellow precipitate with molybdate of ammonia.	Lüneburg
II uble fi		Compare Sphalerite, p. 92.	
Bot	Give with borax a violet bead (manganese.)	when treated with HCl. (See Div. 5, p. 67.)	
		Fuses at 3.—3.5 with bubbling; soluble in dilute hydrochloric acid.	
i		In the closed tube phosphoresces with a	Kjerulfin
	monia a vellow precipi-	Fuses quietly at 5; insoluble in dilute hy- drochloric acid.	Apatite.
	tate (phospho-molybdate of ammonia).	Reacts like apatite, but also gives much water in the closed tube (26 per cent.).	Brushite.
1 1		Same as above. Water=18 per cent.	Isoclasite

TALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture.	Lustre.	Hard- ness,	Sp. Gr.	Fusibility.	Crystalli sation.
i)²(A l, Fe, Mn)As F) ⁹ .	orange-red.	Prismatic.	Vitreous.	5.	4.	2.	v.
³ O ¹¹ +3aq.	Yellow-red.			3.	-	Easily.	Gran.
'As'O ³⁴ +12aq.	Wax-yellow.	Scaly.	Adamantine.		5.8		v.
 D ¹⁴ +12aq.	Lemon-yellow	Tabular.		-	 3.3		v.
°0°+aq.	Violet to honey-yellow.	Distinct.	Vitreous.	3.5	4.34	Easily.	IV.
fg)SO ⁴ +6aq.	Red to yellow- white.	Distinct.	Vitreous.	2.—2.5	1.89	Easily.	rv.
² AlS'O ¹⁶ +24aq.	Colorless to white.		Vitreous.	12.	1.50		I.
¹⁹ +18aq.	Yellow, red- white.		Silky.	1.5—2.	1.7	-	v .
'+7aq.	White.	Prismatic.	Vitreous.	2.—2.5	1.95	Easily.	IV.
gPO4+12aq.	Yellow to brown-white.	Basal.	Vitreous.	2.	1.7	Easily.	I V .
$[g)^{2}B^{2}O^{5}+H^{2}O.$	Gray-white.	Fibrous.	Silky.	3.	3.42	2.	
16.	Yellow to white.	Scaly.	Pearly.	1.	1.48	1.	VI.
B'O'1+6aq.	White.	Foliated.		2.	1.9—2.	Easily.	Fibrous.
6C1°O ³⁰ .	White, gray- green.		Vitreous.	4.5-7.	2.97	2.	L.
0 ¹¹ +3aq.	White-yellow.			3.—4.	8.	Easily.	
B°O ¹¹ +8aq.							
 O ^{\$} +MgF ^{\$} .	Yellow.		Vitreous.	5.5	8.07	8.5	v .
⁻² O ⁸ +CaF ² .	Pale-red.		Greasy.	45.	3.15	8	v.
°O ⁸ +Ca(Cl,F) ² .	Sea-green, blue, yellow, red, white.		Vitreous.	5.	2.9—8.2	4.5-5.	ш.
04+2aq.		Perfect.	Pearly- vitreous.	2,-2.5	2.21	Easily.	v.
)°+5aq.	Snow-white.	Perfect.	Pearly- vitreous.	1.5	2.92	·	v .

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MINERALS WITHOUT METALLIC LUSTRA

B. Fusible from 1-5, and non-volatile.

II. Yield no metal or magnetic mass with sode.

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DIVISION 2 (concluded).

Devestor 8 (in part).

IL MINERALS WIT

	thout		General Characters.	Specific Characters.	Specie
	DIVIBION 2 (Continued.) is in hydrochloric acid without	tinisation.	Fuse at 2, coloring the flame purple-red (lithia). Phosphoresce with a	yellow precipitate with molybdate of am- monia.	Amblygo
	VIBION	gela	light-blue light.	A like mineral, with 4 per cent. of water. Gives a pure lithia flame.	HEBRON
3 m.0.88.	DIT Soluble		With salt of phosphorus in O. F. give a yellow glass which in R. F. becomes	tion in HCI has a yellow color, and gives	Autunite
etie			green (Ü).	Compare Torbernite. Div. 3, p. 75.	
e or magnetic mass.			In matrass gives little wa- ter. B.B. fuses to a clear glass, tinging the flame green.	The dilute acid solution colors turmeric paper red (boric acid).	Datolite
no metallio globule	poration.		The dilute HCl solution gives with sulphuric acid a precipitate of the sul- phate of baryta.	Prismatic cleavage perfect.	Edington
oe no meta	oas wodn hu	water.	Fuses quietly at 2, without swelling or intumescence, to a clear transparent glass.	variationate of ammonia produces note of no	
with soda on charcoal give	Division 8. Soluble in hydrochioric acid, forming a stiff Jelly upon seaporation.	tubegive	Fuses with intumescence. In the HCl solution chlo- ride of barium produces a precipitate. (BaSO ⁴ .)		Ittnerite.
soda on cl	DIVISION 3.	in the closed		Fuses to a voluminous frothy shining slag, which in R.F. further fuses to a vesicular slightly transparent globule; becomes elec- tric on heating.	SCOLECI
with	hloric	B. B. 1	Sometimes curls up in worm-like forms on fu-	Fuses, emitting air-bubbles to a white trans- lucent enamel.	LAUMON
B.B.	pdroc	8)]	sion.	Fuses with difficulty on the edges, worming like scolecite.	Chalcom phite.
H	tn			Resembles scolecite, but is not pyroelectric.	Mesolite.
	nou			Resembles scolecite, but is not pyroelectric.	d a Amblygo HEBRON S Autunite Datolite Edington Edington Ittnerite SCOLECT LAUMON Chalcom phite. Mesolite Thomson
	80		Fuse at 3 with slight in- tumescence.	Often decrepitates. Found in rectangular- terminated crystals. Often in twins con- sisting of 3 or 4 crystals united around a common axis.	Phillipsit
				Usually has the appearance of the square octahedron.	Gismond
			Compare okenite, apophyllite, analoite, be- longing to the next sec- tion.		

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METALLIC LUSTRE.

. 100	Vitreous Vitreous	δ. δ. δ.5	8.11 8.04 8.1 8.1 8.1 8.1 8.1 8.1	2. 2.5 Easily.	VI. VI. IV. V. II.
to sul- rellow. Basal. s, groen, -red. ink. Prisma	Pearly. Vitreous atic. Vitreous	2.—2.5 5.5	3.1 3.1 3.	2.5	IV. V.
s, groen, -red. ink, Prisma	Vitreous atic. Vitreous	5.5	3.	-	▼.
ink. Prisms	atic. Vitreous			Easily.	
ink. Prisms	atic. Vitreous			Easily.	
		4.5	2.7	Easily.	п.
o red.	Vitacana				1
1	Vicreous	. 5.5	2.25	2.	I V .
y.	Vitreous	. 5.5	2.4	Easily.	I.
Prisma	atic. Vitreous	. 5.5	2.2	2.2	v .
ra y , Prisma	atic. Pearly.	8.5	2.3	Easily.	v .
	Glassy.	5.	2.54		111.
Fibrou	18. Silky.	5.	2.3	Easily.	?
Prisma	atic. Vitreous.	5	2.35	2.	IV.
ed).	Vitreous.	. 4.—4.5	2.2	8.	I V .
	Splenden	ıt. 4.5	2.26	Easily.	17.
	ed).	ed). Vitreous	Prismatic. Vitreous. 5. ed). Vitreous. 44.5 white, Splendent 4.5	Prismatic. Vitreous. 5. 2.35 ed). Vitreous. 44.5 2.3 Prismatic. Vitreous. 44.5 2.3	Prismatic. Vitreous. 5. 2.35 2. bd). Vitreous. 4.—4.5 2.2 3. white, Subscience 4.5 2.2 3.

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MINERALS WITHOUT METALLIC LUSTRE

B. Fusible from 1-5, and non-volatile.

1. Field no metal or magnetic mass with soda.

DIVISION 8 (concluded).

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			General Characters.	Specific Characters,	Specie				
			Compare datolate of previous section.		1				
				Heated with hydrochloric acid evolves sul- phuretted hydrogen; not cleavable.	Helvite.				
		ľ	Give with borax in O. F. an amethystine glass (cxide of manganese).	B.B. with soda on charcoal a slight coating of zinc. Heated with HCl evolves H ² S.	Danalite.				
			29 Compare Willemite, p. 91.	Gives off no sulphuretted hydrogen. Per- fect cleavage in one direction.	TEPHRO				
mass.				Color sky-blue. Fuses with difficulty at 4.5 to a white glass.	Hauynite				
retio 1	ion.		With soda on charcoal give	Color sky-blue. Gives off sulphuretted hy- drogen when treated with HCl.	Lapis-La				
magn	aporat		a sulphur reaction.	Fuses quietly at 4.5. Mostly crystallized in rhombic dodecahedrons.	Nosite.				
bule or	aə uodn	or but traces		Fuses at 3 with intumescence. Massive granular.	Scolopsit				
with sola on charcoal give no metallic globule or magnetic mass.	DIVISION 3.—(Continued.) to avid, forming a stiff felly	give no water, or	Fused with a bead of salt of phosphorus which has been saturated with oxide of copper, tinge the flame blue (chloride	In the dilute HCl solution turneric paper assumes an orange color (reaction for zir- conia).	Eudialyt				
al give n	id, formin	ed tube giv	of copper). In the nitric solution nitrate of silver gives a precipitate of the chloride of silver.	Fuses to a clear colorless glass	SODALIT				
a charoe	DIVID Morie ac	b) B. B. in the close	 b In hydrochloric ac b) B. B. in the close 	hydrochlor B. B. in th	b) B. B. in the close	b) B. B. in the close		Fuses with intumescence to a vesicular glass which cannot be perfectly rounded by fusion.	
ith soda on	e in hydroc						b) B. B. in	b) B. B. in	b) B. B. In
B. B. w	Solubl		tion of the silica, ammo- nia gives a precipitate.	Does not give the above reaction with oxalate of ammonia. Found massive and in hexa- gonal prisms. Fuses without intumescence.	NEPHEL (Elacol				
1.1.1	11.23			Tompare Cancrinite, Div. 1, p. 81.					
H.				Behaves like melilite, but is less fusible. $F=4$.	Barsowit (var. Au ite).				
			little or no precipitate,	Fuses quietly to a colorless translucent glass.	WOLLAS				
			but carbonate of ammo- nia causes a copious sep- aration of carbonate of lime.	Compare Pectolite, Div. 4, p. 85.					
				Also compare the difficultly fusible minerals Gehlenite, Div. 5, p. 94; Tachy- lite, Div. 4, p. 86; and Willemite, Div. 2, p. 91.					

METALLIC LUSTRE.

Composition.	Cuice.	Cleavage or Fractara.	Lastre.	Hard-	Bp. Gt.	P	
, Mn, Fe) ² 8iO ⁴ + n,Fe)S.	Wax or honey yellow.		Resinous.	6.—6.5	3.8	3.	L.
Mn,Fe,Zn)*SiO* + >,Mn,Zn)S.	Flesh-red, gray.		Vitreous.	5.5-6.	3.43	Easily.	I
i04.	Reddish- brown, ash- gray.		Vitreous.	6.	4	3.5	IV.
,Ca) <u>AlSi²O⁴+(Na³,</u>)SO ⁴ .	Green to blue.	Dodecahe- dral.	Vitreous.	5.5-6.	2.5	4.5	1
a³, A1, 8, Si.	Azure-blue.		Vitreous.	55.5	24	3	r
AlSi ² O ⁴ +Na ² SO ⁴ .	Gray to black.		Vitreous.	5.5	2.3	4.5	I
⊾Ňa³,Š,Cl,Ši,Ĥ³.	Gravish- white.	Splintery.	Resinous.	5.	2.53	3	Manite
Ca,Fe) ² (Si,Zr) ⁴ O ¹⁴ IaCL	Bose to brown- red.	Basal.	Vitreous.	5.5	29	25	ш.
ElSi ² O ³ +2NaCL	Gray. green, blue. yellow- white.		Vitreous.	5.5—6.	23	3.5_ L	 ב
48i9O24.	Colorless to white.		Vitreous.	5.5	9.7	3	IL.
Ca, Mg) ¹² (A l, Fe) ²)**.	White, yellow, brown.	Basal.	Vitreous.	5.	2.95	 & 	II .
) ^{\$} AlSi ^{\$} O ^{\$} .	Coloriess and green-red.	Hexagonal.	Vitreous to greasy	5.5—6.	2.6	3.5	m.
i ²O⁸.	White.	Granular.	Vitreous.	5.5-6.	2.75	ـــــــــــــــــــــــــــــــــــــ	VL
e <u>.</u>	White-gray.	Basal.	Vitreous.	4.5-5.	8.9	4.5	▼.
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MINERALS WITHOUT METALLIC LUSTRE.

B. Fusible from 1-5, and non-volatile.

II. Yield no metal or magnetic mass with soils.

Deveneour 4 (in part).

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	a perfect jelly.		General Characters.	Specific Characters.	Speci	
			With borax gives the ame- thystine color of man- ganese.	Treated with HOl evolves chlorine, and silica separates as a slimy powder. Gives 9 per cent. of water on ignition.	Klipstein	
			Easily decomposed by HCl,	Fuses with slight intumescence to a white enamel-like glass. Yields but little water. After fusion gelatinizes perfectly with hydrochloric acid.	PECTOLI	
			the separation of the	but slightly attacked by acids.	Арорну	
magnetic mass.		"Tech Jent."	silica, the solution gives with ammonia no or only a slight precipitate.	 Fuses at 2.5—3, with frothing to a milk-white glass; yields much water; after fusion but slightly attacked by acids. Compare Xonaltite and Sepiolite, Div. 5, p. 93. 	Okenite.	
ground or mag	without forming a p		Decomposed by HCl like the preceding. After the separation of the silica the solution gives with ammonia a copious precipitate.	B. B. at first becomes opaque, but fuses quietly to a clear glass. Occurs usually in trapezohedrons and cubes.	ANALCII	
nonemose.	silica with	the closed tube give water.	The dilute HCl solution gives with sulphuric acid a white precipitate (BaSO ⁴).	Fuses at 3 with intumescence. (Contains 13 p. c. of water.) Compare Harmotome, p. 87.	Brewster	
-	DIVISION 4. a residue of silica	ad tube	and fuse with contor- tions to enamel - like masses. In the solu- tion from which the silica has been separated ammonia produces a pre- cipitate.		Prehnite	
and m	DIVI Soluble in hydrochloric acid, leaving a res	he close		Distinguished by its rhombohedral crystalli- zation and imperfect cleavage.	Снавалі	
soda on charcoal give no metallic		B. in t		Perfectly cleavable in one direction. Ortho- rhombic. B. B. intumesces strongly.	STILBITI	
				Perfectly cleavable in one direction. Mono- clinic. Lustre very pearly on one face. B. B. intumesces strongly.	HEULAN	
	Mor			One perfect cleavage. Intumescence less.	Hypostill	
man	udroc		1	Fuses with scarcely any intumescence.	Mordenit	
i	ie in A				Fuses at 3.5—4 with intumescence; not cleavable. (Water = 9 p. c.)	Chonicrit
IL B.	Solub		These minerals, the hard- ness of which is not above 3, are softer than,	Fuses quietly at 4.; cleavable in one direc- tion. (Water = 11 p. c.)	Pyroscler	
		, 4445 1		Exfoliates in worm-like forms.	Vermicul	
				Exfoliates prodigiously.	JEFFERI	
				Swells up; fuses with difficulty. (Water = 13 p. c.)	Jollyte.	
			the other minerals of this division.	Swells up and fuses to a white enamel. (Water = 21 p. c.)	Kerrite.	
				Swells up and fuses to a brown glass. (Water = 11 p. c.)	Maconite	
				Fuses with difficulty to a white enamel. Water = 4 p. c.)	Willcoxit	
				Exfoliates slightly; fuses with difficulty to a brown-yellow blebby mass. (Water = 13 p. c.)	Dudleyit	

ETALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture.	Lastre.	Hard- ness,	Sp. Gr.	Fusibility	Crystalli- sation.
ı,Ŝi, Ĥ ².	Dark liver- brown to black.		Dull to sub- metallic.	5-5.5.	8.5		Amorph
\ ²Si³O⁰.	White to gray.	Fibrous.	Silky.	5.	8.7	2	v.
.Si ² O ⁶ + aq) +	Colorless, white, rose red, yellow.	Basal.	Vitreous pearly.	5.	2.3	1.5	п.
i ² O ⁶ +aq.	White.	Fibrous.	Pearly.	4.5-5.	2.8	Easily.	IV. ?
3i4O19+2aq.	Colorless to white, gray, green, yel- low, red.	Not cleav- able.	Vitreous.	5.—5.5	2.28	2.5	I.
±lSi ⁶ O ¹⁶ +5aq.	Yellowish- white to gray.	Prismatic.	Pearly vit- reous.	5.	2.45	8.	v .
rlSi ³ O ¹⁹ .	Apple to oil green, white.	Basal.	Vitreous.	66.5	2.9	2.	IV.
CaAlSi ⁵ O ¹⁵ +6aq.	White, flesh- red.		Vitreous.	45.	2.1	Easily.	111.
^{\$})AlSi ⁶ O ¹⁶ +6aq.	White, yellow- red.	Prismatic.	Pearly vit- reous.	8.5-4.	2.16	22.5	IV.
⁶ O ¹⁶ +5aq.	White-red.	Clinodiag- onal.	Pearly vit- reous.	8.5-4.	2.2	22.5	v .
2)2Al2Si9O26+12aq	White.	Fibrous.	Vitreous.	8.5-4.	2.2	Easily.	
')AlSi'0''+6aq.		Concretion- ary.	Silky.	5.	2.08	Easily.	
)10AlSi7O30+6aq	White-yellow.		Sil ky.	2.5-8.	2.9	3.5-4.	
'Si'0 ^{**} +12aq.	Apple to em- erald-green.	Micaceous.	Pearl y .	8.	8.74	4.	V. ?
12 Al Si O36 + 12ag	Brown-yellow.	Micaceous.	Pearly.	1.5	2.75		VL ?
Fe)'Si'O*0+6aq.	Brown-yellow.	Micaceous.	Pearly.	1.5	2.8		IV. ?
) ⁶ Al'Si ⁹ O ³⁶ +12aq				8.	2.61	Difficult.	Amorph.
3i*O19+10aq.	Greenish - yel- low.	Micaceous.	Pearly.	1.5	2.3		
,Fe)*Si4O17+5aq	Dark-brown.	Micaceous.	Pearly.	2.	2.8		-- - ------------- - --------- - ---------------------------- - ---
,Na ²) ⁶	Gray.	Micaceous.	Pearly.	1.5			
Si ¹ O ²⁹ +10ag.	Bronze.	Micaceous.	Pearly.		· ·		

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MINERALS WITHOUT METALLIC LUSTRE

B. Fusible from 1-5, and non-volatile.

II. Yield no metal or magnetic mass with so?

DIVISION 4 (concinded).

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II. MINERALS WJ

		ct jelly. ucd) B. B. in the closed tube give water.	General Characters,	Specific Characters,	Species										
				Fuses quietly at 3. to a milk-white globule. The dilute HCl solution colors turmeric paper orange-yellow (zirconia).	Catapleiit										
			the closed	the closed	the closed	the closed	the closed	the closed	the closed	the closed	$\mathbf{H} = 44.5. \text{Cleavable.}$	Fuses at first with intumescence, then quietly at 2.5-3. to a yellow-brown glass. With salt of phosphorus in the reducing flame gives a violet color (titanic acid).	Mosandrit		
88				Absorbs water with avidity. ($\dot{H} = 10 \text{ p. c.}$)	SEPIOLIT										
II. B. B. with sola on charood give no metallic globule or magnetic mass.	it jell		Difficultly fusible (F=5.)	Does not absorb water. (Water = 20 p. c.)	DEWEYLI										
	ning a perf	a) (Contin	Fuses at 2.5 to an opaque black shining glass. Dif- ficultly decomposed by hydrochloric acid.												
	ut for	F		Compare Peotolite, Choniorite and Preh- nite of the preceding subdivision.											
	()		Div. 3, p. 84.												
	Drvision 4.— (Continued.) eaving a residue of silica u	estance of stitue	Micaceous; also scaly mas- sive.	Fuses easily in the candle flame, and B. B., with intumescence to a gray enamel giv- ing a lithia-flame.	Cryophyll										
	4(0		Fuses easily to a black slag.	The silica separates as gelatinous lumps.	Tachylite										
	NOISI	but traces.	color (titapic acid)	Fuses quietly. Difficultly decomposed ; the silica separates as a slimy powder.	Schorlom										
	Drv ochloric acid, leavi	 b) B. B. in the closed tube give no water, or but traces. 			Tscheffkir										
			(Columbium reaction).	÷.	4	÷ ÷	÷.	4	÷ ÷	give no	2.5 to a white vesicular	Cleavable in two directions.	Werner (Scapoli		
	ndr													ly be further fused.	Occurs in glassy crystals.
	Soluble in 1									flocks; the acid solution when boiled with tin be- comes beautifully blue		100			
				Cleaves in two directions	Fusibility = 3.5. Often striated, and shows beautiful play of colors.	LABRADO									
				Fusibility = 4.5 . Gelatinizes with acids.	Anorthite										
			Gives the chlorine reaction with oxide of copper.	Difficultly fusible.	Microsom										
			Sphene and Danburite, Div. 6, p. 87; also Teph- roite, Div. 3, p. 84.												

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METALLIC LUSTRE.

Jomposition.	Color.	Cleavage or Fracture.	Lustre.	Hard-	Bp. Gr.	Pusibility.	Orystalliza tion.
ı)(Si,Zr)'O ⁹ +2aq	Yellow-brown.	Prismatic.	Vitreous.	6.	2.8	8.	ш
Ďi,Ċa,Ňa²,Ťi,Ŝi,	Reddish- brown,	Prismatio.	Resinous.	4	2.95	8.	IV. ?
$D^3+2aq.$	Gray-white.		Dull.	22.5		5.	
D ¹⁰ +5aq.	White, yellow, red.		Greasy.	3.	2.2	5.	
мg ,Ŝi, Ĥ .	Gray-black.		Resinous.	2.5	2.58	2.5	
	·			-			
Fe³ (A 1, F e)'Si ²¹	Black, green to brown-red.	Micaceous.	Pearly.	2.5	2.91	1.5-2.	IV.
Ċa, Ňa², Äl, Ši, Ĥ²	Gray, pitch- black.		Vitreous.	6.5	2.6	2.5	
8i,Ti) ¹² O ³⁹ .	Black.		Vitreous.	77.5	3.8	3.	
Ĵa,Ťi,Ŝi.	Black.		Vitreous.	5.—5.5	4.5	Easily.	
² ,K ²)AlSi ² O ⁸ .	White, gray, blue, green, red.	Prismatic.	Vitreous to greasy.	56.	2.6—2.8	2.5	п.
Si ⁹ O ⁸⁶ .	Colorless,White	·	Vitreous.	5.5	2.7	3.	п.
Ši,Žr,Ĉbª.	Yellow-brown.	Prismatio.	Vitreous.	5.5	8.41	8.	IV.
') A 1Si ⁸ O ¹⁰ .	White, gray- brown, green.	Angle 94°.	Vitreous.	6.	2.7	8.5	VI.
² O ⁸ .	Colorless, white, gray.	Two equal cleavages.	Vitreous.	6.—7.	2.7	4.5	VL.
AlSi ² O ⁸ , CaSO ₄ ,	Colorless.		Vitreous.	6.	2.6	5.	ш.

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MINERALS WITHOUT METALLIC LUSTRE.

B. Fusible from 1-5, and non-volatile.

II. Field no metal or magnetic mass with sodis.

DIVISION 5.

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Division 6 (in part).

	olor an	General Characters.	Specific Characters.	Speci
	the co	Gives water in the matrass	s. Found in fibrous radiated masses.	Carpholi
	tox B. y hydrochloric acid. p amelhystine color id (oxids of man	Fuses quietly at 3. Cleave	age indistinct, sometimes dodecahedral.	SPESSA (manga garne
	DIVISION attacked by hi give a deep an boram bead	Fuses with intumescence a	at 2.—2.5. Plainly cleavable in one direction.	Piedmon
	Blightly atta B. B. give to the bor ganese).	Fuses quietly at 3. Plain	ly cleavable at an angle of 92°.	RHODO (mang span
	Compare Azinite in n	ext section.		
magnetic mass.		Fused alone colors the flame green (boric acid).	Fuses at 3 to a globule which while hot is clear, but becomes cloudy on cooling. Yields no water in the closed tube.	Danburi
reti		minto Broom (source mondy.	Yields water in the closed tube.	Howlite
5		The powder is soluble in hydrochloric acid, leav- ing a yellow residue of tungstic acid.	but soon bleaches on dilution.	Scheelit
globule		Micaceous. Give to the	Fuses at 2. Gives in the closed tube little or no water.	Lepidol
allie .		blowpipe flame the pur- ple-red color of lithia.	B. B. vermicular exfoliations. Gives in the closed tube much water.	Cookeite
give no metallio		Micaceons, but do not give	Swells up B. B. and gives much water in the closed tube (11 per cent.)	Thermoj lite (s tine).
	dona.		Fuses quietly. Easily decomposed by sul- phuric acid.	Euphylli
charcoal	divis	the lithia liame.	Fuses quietly. Difficultly decomposed by sulphuric acid.	MARGAR
no	c 6. regoin		Compare Muscovite and Biotite, Div. 6, page 94.	
soda	DIVISION 6.	Not micaceous. Give to	Fases quietly to a white enamel. (B) Com- pare Amblygonite, p. 83.	PETALIT
with	D1 Ing to	purpre-red color or minus.	Intumesces, throwing out fine branches, which fuse to a clear glass.	SPODUM
. B. B.	DIVISION 6. A belonging to the foregoing	Phosphoresces when heat- ed, or when struck with a hammer.	Fuses quietly at 3 to a transparent colorless glass. With salt of phosphorus in the open tube gives the fluorine reaction.	Leucoph
H.	Not	Gives water in a matrass; the partial HCl solution gives a precipitate with sulphuric acid (baryta).		HARMOT
		Fused with a mixture of fluor-spar and bisulphate of potassa momentarily	glass. The fine powder of the fused min- eral gelatinizes with acids.	AXINITE
		colors the blowpipe flame green.	Different varieties vary much in blowpipe characters; all become electric by heat- ing. Some gelatinize after fusion.	Tourma
		Partially decomposed by hydrochloric acid; the	Fuses with slight intumescence to a black-	Titanite (Sphene)
	c < 0	solutions when boiled	Differs in crystalline form.	Guarinit
1	1	with tin become violet (titanic acid).	Fuses with brisk intumescence to a blackish mass.	Keilhaui (Yttrotita

'ALLIC LUSTRE.

mposition.	Color.	Cleavage or Fracture.	Lustre.	Hard- ness.	Sp. Gr.	Fusibility.	Crystalli- zation.
1, F e,Mn)Si ² O ¹⁰ .	Straw-yellow.	Stellate, fibrous.	Silky.	5.—5.5	2.9	8.5	IV.
⁵AlSi³O¹⁵.	Brownish-red.		Vitreous.	7.	4.2	8.	L
n,Fe,Al) ⁸ Si ⁶ O ²⁶	Cherry-red to reddish-brown	Prismatic.	Vitreous.	6.5	3.4	8.	v .
	Rose-red, brown.	Prismatic.	Vitreous.	6.	3.6	2.5	VI.
) ⁸ .	Pale-yellow.		Vitreous.	7.	2.9	3.	VI.
0 ²³ +5aq.	White.		Sub-vitreous.	3.5	2.55	Easily.	Amorph.
	White, brown, green-red.		Vitreous.	4.5—5.	6.	5	п.
14Si19O89.	White-gray pink.	Micaceous.	Pearly.	2.5	3.	2.5	īv.
1,Ŝi,Ĥº.	White.	Micaceous.	Pearly.	2.5	2.7	Diffi- cultly.	
¹ +2aq.	Brown to white.	Foliated.	Pearly.	2.5	2.6	5.	
K ⁶) ⁶ Si ⁹ O ³⁶ +4aq	White.	Foliated.	Pearly.	3.5	2.8	44.5	
Si ² O ¹² .	White, red, gray.	Micaceous.	Pearly.	4	2.99	44.5	
\$i ⁶ O ¹⁶ .	White, gray- pink.	Basal.	Greasy.	6.5	2.45	3.5	.
i ³ O ⁹ .	White-gray, green-pink.	Prismatic.	Pearly.	6.5	3.18	3.5	₹.
(Ca,Be)4Si3O10.	Green-white.	Basal.	Vitreous.	8.5-4.	2.97	8.	IV.
)14+5aq.	White-red.		Vitreous.	4.5	2.45	8.5	IV.
K²) ¹ (Al, Fe,B) ⁸	Clove-brown to pearl-gray.		Vitreous.	6.5-7.5	8.27	2.	VL
<u>.</u> Mn,Mg,K²,Ňa², F.	pink. white.		Vitreous.	6.57.5	 2.9	8.—5.	 III.
·.	Brown, green, yellow, black.	Prismatic.	Vitreous.	55.5	8.5	8.	v .
\$	Honey-yellow.	Prismatic.	Vitreous.	6.	3.48	8.	II.
, Ä l,Ši,Ťi.	Brown black.		Resinous.	6.5	3.7	8.	IV.

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MINERALS WITHOOT METALLIC LUSTRE

B. Fusible from 1-5, and non-volatile.

II. Field no metal or magnetic mass with sods.

DIVERION 6 (concluded).

II. MINERALS WITH

1		General Characters.	Specific Characters.	Specie
			Fuses at 5. Has two perfect cleavages at 90°.	Orthocla
			Fuses at 4. Shows strictions on one cleav- age surface.	Albite.
		Hardness 6. Fuse quietly.	Fuses at 3.5. Striations as above.	Oligoclas
			Fuses at 3.5. Striations as above. Gives water in the closed tube. Phosphorescent.	Tscherma
.0.88.			Fused with soda, the silica separated from the hydrochloric solution, gives with sul- phuric acid a precipitate (baryta).	Hyalopha
0 11			Compare Labradorite, Div. 4, p. 86.	la serie de la ser
gnet		Hardness=6.5. Fuse with	Fuses to a white or yellow slag.	ZOISITE.
ule or ma		swelling and intumes- cence to a slaggy mass. They gelatinize with acids after fusion.	Fuses to a black or dark-brown slag.	Epidote tacite).
etallic glob inued.)	(approx	Hardness, 6.5—7.5. Gela- tinize with acids after fusion.	Fuses quietly at 3 (grossular) to 4.5 (pyrope	
	inued.		Fuses with intumescence at 3.	Vesuvia (Idocra
nom	(Cont		Resembles grossular (but does not gelatinize after fusion).	Monzoni
B. with soda on charcoal give no metallic globule or magnetic mas .	DIVISION 6 (Continued.) Not belonging to the foregoing divisions.	Hardness 6. Cleavable at an angle of 93°.	Includes many varieties, from the colorless diopside and white malacolite to black augule; light-colored varieties fuse to a white glass, while the dark give a black glass. The species is recognized by the cleavage and habit of crystal, the variety only by experience.	Pyrozen
soda			Fuses to a white glass.	Tremolit
ith		Service and the service of the	Finely fibrous with fibres easily separable.	Asbestus
B. w		Hardness 5.5. Cleavable at an angle of 124°.		Actinolit
II. B. B. with sods on charcoal give no metallic globule or magnetic mass.			As above under pyroxens. The species in- cludes tremolits, asbestus, actinolits, and many darker colored varieties. Can be recognized by the cleavage, but the varie- ties can only be learned by experience.	(Hornhla
		Fuses at 4. Exfoliates, and yields water in a matrass.	Occurs in thin short fibrous layers.	Gümbeli
		Fuses at 2. Gives water in a matrass.	Fuses with intumescence to a white glass.	Wilson (alt'd S lite).
		Fuse with swelling up, at	Characterized by an intense vitreous lustre.	OBSIDIA
1		3.5-4. to a vesicular	Characterized by a strong fatty lustre.	PITCHST
		white glass or enamel. They are amorphous,	Characterized by a mother-of-pearl lustre;	PEARLST
		volcanic products, and	sometimes yields water.	- Dicheot

ETALLIC LUSTRE.

omposition.	Color.	Cleavage or Fracture.	Lustre,	Hard- ness,	Sp. Gr.	Fusibility.	Orystalliza tion.
5O1 6 .	Colorless, white, flesh- red, gray- green.	Right-angle.	Vitreous.	6.	2.5—2.6	5.	v.
i ⁶ O ¹⁶ .	Colorless, white-gray, dull-green.	93°30′.	Vitreous.	6.	2.6	4.	VI.
,K²) AlSi⁵O¹4.	White, flesh- red.	93°.	Vitreous.	6.	2.6-2.7	3.5	VI.
ı) A lSi⁵O¹4.	Gray-white.	94°.	Vitreous.	6.	2.64	8.5	VI.
AlSi ⁴ O ¹⁹ .	White, flesh- red.	2 cleavages.	Vitreous.	6.	2.9	5.	v .
±1 F e) ³ Si ⁶ O ²⁶ .	White-ash gray.	Prismatic.	Vitreous.	6.—6.5	3.—8.3	33.5	IV.
11Fe) ³Si⁵O²⁵.	Gray, pista- chio-green, brown-yel- low.	2 cleavages.	Vitreous.	6.—7.	8.2-8.5	8.—3.5	٧.
$\overline{\mathcal{D}^{19}}$. $\mathbf{R} = \Lambda \mathbf{l}, \mathbf{F} \mathbf{e}, \mathbf{C} \mathbf{r}$ a, Mg , Fe, Mn.	White, red- brown, black.	Dodecahe- dral.	Vitreous.	6.5-7.5	3.2-4.3	34.5	I.
Fe) ² Si ⁷ O ²⁸ .	Brown-green, yellow, blue.		Vitreous.	6.5	8.8-3.4	3.	11.
Ĵa, Ňa², Ŝi.	Gray-green.		Vitreous.	6.	8.	8.	Mass.
s,Mg,Fe,Zn,Mn, a²(±1,Fe, M n).	Colorless, white, gray, brown, green, and black.	87° & 93°.	Vitreous.	5.5—6.	8. 2— 8.5	2.5- 5.	v .
)SiO ⁸ .	White.	Bladed.	Pearly, vitre- ous.	5.5	2.9-3.1	8.5	v .
,Fe)SiO ⁸ .	White.	Fibrous.	Sil ky .				
,Fe)Si Q ⁸ .	Green, brown.	124°30′ and 55°30.	Vitreous.	5.5	8.—3.2	4.	₹.
,Mg,Fe,Mn,Na ³ , ³ (Al,Fe,Mn).	Like pyroxene	124°30 and 55°30'.	Vitreous.	5.5	2.9—8.4	2.5- -5.	♥.
я́, н .	Green-white.	Fibrous.	Pearly.			4.	
Йg,Ŝi,Ĥ¹.	White to red.	Cleaves at right angles.	Dall.	8	2.7	2	II.
≿, Mg,Ŕ²,Ňa²,Ŝi.	White, gray, green, yellow, black.	Break with sharp edges. Conchoidal.		6.	2.2—2.8	8.1-4.	

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MINERALS WITHOUT METALLIC LUSTRE

O. Infusible or fusible above 5.

Alevanour 1 (in past).

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		General Characters.	Specific Characters.	Spe		
			Insoluble in hydrochloric acid. (13 p. c. water.)	Alunit		
		With soda on coal give a sul-	Easily soluble in hydrochloric acid. (47 p. c. water.)	ALUM		
	Ì	phur reaction,	B. B. becomes black, burns and falls to pieces.	Pissop		
1 1		Like aluminite. (37 p. c. water.)	Felsob			
		1	Compare Kalinite, Tschermigite, and Alunogen, which are soluble in water.	-		
color.		With soda on coal gives a glo- bule of lead.	B. B. puffs up and half melts without becom- ing fluid. The solution gives with molyb- date of ammonia a yellow precipitate.	Plum		
DIVISION 1. then molatened with coball solution and again ignited assume a beautiful blue color. (It is recessary to putterize hard anhydrous minerals before fractment.)		With soda on coal gives a zinc coating.	Soluble in hydrochloric acid, forming a per- fect jelly.	Cala (electr min		
beautif		Contains 27 p. c. of water.	WAVE			
ada			Contains 27 p. c. of water.			
assum	1	Give the phosphoric acid reac-	Contains 24 p. c. of water.	Pegan		
Ited	1 H	tion when fused with magne-		Fische		
40m	water.	sium in the closed tube (see 134). Mostly soluble in caustic po- tassa, also if to this solution an excess of nitric acid is added, and some molybdate of ammonia, a yellow preci- pitate is thrown down.	Contains 4 p. c. of water.	Berlin		
again Irous 1	tube give		on Contains 21 p. c. of water.			
and and						
DIVIBION solution c hard an	Deed					
DIVI	the closed		Contains 23 p. c. of water.	Redon		
balt	th th		Contains 12 p. c. of water.	Amphi		
A co	A		Contains 12 p. c. of water.	Tavist		
ued with	a) B.		Contains 21 p. c. of water.	Coert		
molsten			Gelatinizes perfectly. $H = 3$. (Water 42 p. c.)	ALLOP		
B. then (It u		Soluble in hydrochloric acid,	Has a lamellar structure. $H = 4$. (Water 30 p. c.)	Samoi		
Påret ignited B.		with the separation of gela- tinous silica.	Very soft. $H = 1-2$. (Water = 16 p. c.)	Halloy		
Pitrat (g			Very soft. $H = 1-2$. (Water = $33\frac{1}{2}$ p. c.)	Collyri		
		Fused in a closed tube with bi- sulphate of potassa gives the fluorine reaction.	Gives water in the closed tube, which reacts for fluorine.	Ralsto		
		Easily soluble in caustic potassa.	Hardness = 2.5-3. Water 341 p. c.	GIBBS		
			Water = 15 p. o. H = 6.5.	DIASP		
		Very easily cleavable in one di- rection.	Water = 13 p. c. H = 12.5.	Kaolir		
			Water = 15 p. c. $H = 1$. Occurs in scales.	Pholer		

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METALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture.	Lustre.	Hard- ness,	Sp. Gr.	Crystalliza- tion.
1°S40?2+62q.	White-gray.	Basal.	Vitreous.	8.5-4.	2.6	IIL
)6+9aq.	White.		Dull.	12.	1.66	
e,Ŝ,Ĥ².	Light to olive-green.		Vitreous.	1.5	1.96	
0 ⁹ +10aq.	White-yellow.	Perfect.	Pearly.	1.5	2.33	IV.
ù,°°,Ĥ°.	White, reddish-yellow, gray-green.		Resinous.	4.—5.	4.8	
i 0 ⁴ + H ² O.	Colorless, white-yel- low, green, blue.	Prismatic.	Vitreous.	4.5-5.	8.5	IV.
4019+12aq.(F).	White yellow, gray- brown, blue, green.	Radiated.	Pearly.	3.5-4.	2.3	IV.
⁹ O ¹⁴ +18aq.	White.		Vitreous.	3.5-4.	1.94	
2011+6aq.	Deep - green, gray, white.	1.000	Vitreous,	8,3.5	2.5	IV.
"O11+8aq.	Grass to olive-green.		Vitreous.	5.	2.46	IV.
208+aq.	White-gray-red.		Vitreous.	6.	2.64	Massive.
05+6aq.	Green, yellow, gray- white.		Vitreous.	5.	2.37	Compact
6027+3aq.	Pale-green.		Vitreous.	5.5	3.10	Compact
"O ²⁵ +16aq.	Gray-red.	1.1	Vitreous.	4.	2.53	
e.P2,H2.	Gray-yellow.		Dull.	8.5	2.	Massive.
a, P2, H2.	Milk-white.			6.		Massive.
1P2O11+8aq.	White.	1	Pearly.	1		Acicular.
4019+10aq.	Milk-white to blue.	Uneven.	Vitreous.	5.	2.5	
O ⁵ +5aq.	White, blue, yellow, green.	Brittle.	Resinous, vitreous.	8.	1.87	Amorph.
1ºO12+10aq.	White, gray, yellow.	1	Resinous.	4.5	1.8	Stalac.
'O [†] +4aq.	White, gray, green, yellow, red.		Waxy.	1.—2.	2.	
iO ⁸ +9aq.	White.		Glimmering.	1.—2.	2.1	Amorph.
g,Ña ⁹ ,F,Ĥ ⁹ .	Colorless-white.		Vitreous.	4.5	2.5	L
04.	White, yellow, red.		Dull.	2.5-8.	2.8	111.
04.	White, gray, brown, blue, green.	1	Vitreous, pearly.	6.5—7.	8.4	rv.
O ^T +2aq.	White, gray, brown.		Pearly.	12.	2.5	IV.
3019+4aq.	White, gray, red.		Pearly.	12.	2.5	IV.

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MINERALS WITHOUT METALLIC LUSTRE

C. Infusible or fusible above b.

DEVENDOR 1 (concluded).

water	General Characters.	Specific Characters.	Spec
a live		Tough; can be cut into chips; imperfectly decomposed by sulphuric acid.	Cimolit
closed tube	chalky.	Unctuous. Forms a pasty mass with water.	Clay.
in the d	Pholerite, above; Kaolinite and Pholerite, above; Kaolinit forms the basis of most clays	Water = 35 p. c. Falls to pieces in water.	Schrött
B.B.1	the class of the state of	Water = 25 p. c. Falls to pieces in water.	Milosch
a) (Continued.)		-	
	With soda on coal give a sul	The partial nitric solution gives a reaction for phosphoric acid with molybdate of ammonia.	Svanber
	phur reaction.	Gives no phosphoric acid.	Alumia
	Colors the flame green when moistened with sulphuric acid and ignited.	B. B. swells, loses its blue color, and falls into small pieces. Not acted upon by acids.	LAZULI
	With soda on coal gives a zine coating.	^C Gelatinizes perfectly with hydrochloric acid.	Willem
COR		The micaceous variety swells up B. B. into fan-like forms. Compact or slaty varieties do not exfoliate.	Pyroph
but traces.	Very soft. H = 13.	Unaltered B. B.; unacted upon by acids.	Agalma
or bu		Somewhat decomposed by acids.	Myelin.
water.		Like pyrophyllite.	Westani
18		The foliæ are very elastic. Not acted upon by sulphuric acid.	Muscov
be gives	Very distinctly foliated.	Not so cleavable; foliæ not elastic; decom- posed by sulphuric acid.	SEYBER
sed tul		Fused in the open tube with salt of phes- phorus gives the fluorine reaction.	TOPAZ.
in the closed tube	11 - 40	Fused with a mixture of bisulphate of po- tassa and fluor spar gives a green flame (boric acid). Pyroelectric.	RUBEL: (var. to line
b) B. B. i		Decomposed in a bead of salt of phosphorus leaving a skeleton of silica. Cleavable in two directions at 91 ¹ / ₂ °.	Andalu
0	Not affected by acids.	Decomposed like the preceding. In bladed	Cyanit
1		Commonly fibrous. Decomposed like the preceding.	FIBROL
		Slowly but perfectly soluble in salt of phos- phorus; very hard; $H = 9$.	Corund
		Slowly but perfectly soluble in salt of phosphorus. $H = 8.5$.	CHRY BER
C Period to ha		which is not above 6, and Cassiterite, which	-

IETALLIC LUSTRE

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Composition.	Color.	Cleavage or Fracture,	Lustre.	Hard- ness.	Sp. Gr.	Crystallise tion.
1º0º4+6aq.	White, gray, red.		Dall.	Soft.	2.2	Amorph,
i,Ĥ².	All colors.		Dull.	Soft.	1	Amorph.
i ⁸ O ⁸⁰ +30aq.	White-green.		Resinous.	3.5	2.	
∃r)SiO ⁵ +3aq.	Blue-green.			1.5	2.13	
a,Ňa ¹ ,P ² ,Ī,Ĥ ³ .	Yellow, or yellowish- brown		Vitreous,	5.	8.3	ш.
09.	White.		Vitreous.	23.	2.74	?
Fe) #1P2O9+H20.	Azure-blue.		Vitreous,	5.—6.	3.1	v.
iiO4.	Colorless, brown, yel- low, red, green.		Vitreo-re- sinous.	5.5	8.9-4.2	ш.
⁸ O ⁹ +H ² O,	White, gray, green.	Micaceous to scaly.	Pearly.	12.	2.9	IV.
7²,Ši,Ĥ².	White, gray, green.	Massive.	Dull.	22.5	2.8	Massiye.
0 ^s .	Yellow-red, white.		Dull.	2.	2.5	
i,Ĥ ⁹ .	Brick-red.		1.000	2.5	1111	Radiated
1SiºO ⁸ .	Gray, white, brown, green.	Micaceous,	Pearly.	2.5	2.8-3.	IV.
,Ca ⁶ , Al ⁹ , Fe ⁹)SiO'.		Foliated.	Pearly.	4.5	3.1	IV.
(0, F ²) ⁵ .	Colorless, white, blue, green, yellow.	Basal.	Vitreous.	8.	3.5	IV.
(a, K) ⁶ Al ⁶ B ³ Si ⁶ O ⁴⁵ .	Violet, rose-red.		Vitreoua.	7.5	8.	ш
05.	White, gray, yellow- red.	Prismatic.	Vitreous.	7.5	3.2	IV.
O ⁸ .	KANGLA DISEAS	Prismatic.	Vitreous, pearly.	57.	8.6	VL.
O ⁸ .	White, brown, green, red.	Prismatic,	Vitreous.	67.	3.2	v .
1	White, gray, blue, all colors.	Rhombohe- drai.	Vitreous.	9.	4.	III,
104.	Aspazagus to emerald- green,		Vitreoua.	8.5	8.7	IV.

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(**1**°age 91)

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MINERALS WITHOUT METALLIC LUSTRE

C. Infusible or fusible above b.

DIVISION 9

DIVISION 8

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IL MINERALS WITH

]	General Characters.	Specific Characters.	Specie				
	on and la a color.	Dissolve in hydrochloric acid	Gives much water in the closed tube.	Hydro: CITE				
DIVISION 9.	with cobalt solution and B. assume a green color.	with evolution of carbonic acid.	Gives little or no water in the closed tube.	Smithsor				
Ä		With hydrochloric acid form a	Gives much water in the closed tube. Pyroelectric,	Calamin				
	Kolatanad B.	perfect jelly.	Gives no water in the closed tube.	Willemi				
		Compare Goslarite, Spha- lerite, and Cassiterite.						
			Dissolves easily and quietly in hydrochloric acid.	Brucite.				
	DIVISION &. After lynition B. B. Acres an alkaline reaction and change the color of moletened turmerto paper to red brown.		As above, gives a strong manganese reaction with borax.	Pyrochroi				
d turmark		oletened turmer	d turmar	od turma	d turmar	Give much water in the closed tube.	Effervesces in hot HCl; the concentrated solution gives no precipitate with sulphuric acid.	Hydron NESIT
					Hydrod mite			
r of mote		Effervesce in hot HCl; the concentrated solutions give a precipitate with sulphuric acid.	Predazzit					
	e color			Pencatite				
	the th	Effervesce and are soluble in cold dilute acid; the dilute	cleavage.	Calcite.				
øð	md ch	solution gives no precipitate with sulphuric acid; but the strong solution does.	Is not cleavable. B. B. falls to pieces.	Aragonit				
KOI	non o		Compare Strontianite, below.					
DIVISION 8.	le reacti aper lo		The concentrated solution gives with sul- phuric acid a precipitate of sulphate of lime.	Dolomite				
	n albalt P	Effervesce and are soluble in hot but not in cold dilute hydrochloric acid.		Magnesi				
	B. Aave a		Compare Siderite and Diallogite, Div. 4, p. 92. Compare also the two following minerals.					
	attion B.	dilute hydrochloric acid; the		Strontiar				
	Var (g	very dilute solution gives a precipitate with sulphuric acid.	Imparts to the flame a yellowish-green color.	Barytoca				
	ľ	Compare <i>Yttrocerite</i> , also <i>Talo</i> and <i>Muscovite</i> , which sometimes have an alkaline reaction after fusion.						

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ETALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture,	Lustre.	Hard- ness.	Sp. Gr.	Orystallin tion.
D⁵ +2a q.	White, gray, yellow.		Dull.	2.—2.5	8.7	
) ³ .	White, gray, green, blue, yellow, red.		Vitreous.	5.	4.4	пі
O⁴+H³O.	White, gray, green, blue, yellow, red.	Prismatic.	Vitreous.	5.	8.5	IV.
O ⁴ (often with).	White, gray, brown, green, yellow, red.		Vitreo-resin- ous.	5.5	4.	III.
·O ³ .	White, gray, green.	 Basal.	Pearly.	2.5	 2.85	 III.
n,Mg)O ² .	White to bronze.	Basal.	Pearly.	2.5		
³ O ¹⁰ +4aq.	White.		Silk y -dull.	8.5	2.1	v .
$\mathbf{M}_{\mathbf{g}}\mathbf{)}\mathrm{CO}^{3}+\mathbf{H}^{2}\mathbf{O}.$	Yellow, gray, green, white.		Vitreous.		2.5	
$0^{3} + H^{2}MgO^{3}$.	White to gray-white.		Vitreous.	8.5	2.63	
°+H°MgO°.	Blue-gray.		Vitreous.	8.	2.5	
8,	Colorless, white, and of all tints.	Rhombohe- dral.	Vitreous.	3.	2.6-2.8	III.
³ .	Colorless, white, yel- low, red, blue.		Vitreous.	3.5-4.	2.9—3.	IV.
[g)CO ^{\$} .	White, gray, brown, etc.	Rhombohe- dral.	Vitreous to pearly.	3.5-4.	2.8-2.9	ш.
)*.	White, yellow, gray, brown, green.	Rhombohe- dral.	Vitreous.	8.54.5	8.—8.1	ш.
۹.	White, gray, yellow, green.	Prismatic.	Vitreous.	8. 5—4 .	8.7	IV.
a)CO ^s .	White, gray, yellow, green.	Prismatic.	Vitreous.	4.	8.6	v .

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MINERALS WITHOUT METALLIC LUSTRE.

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O. Infusible or fusible above 5

DEVENION 4 (in part).

II. MINERALS WITH

Γ	General Characters.	Specific Characters.	Врек
	Colors the fiame carmine-red (lithia).	With salt of phosphorus gives reactions for copper and cobalt.	Lithiop
1	With soda on coal easily re-	Yields little or no water in the closed tube.	Cervant
	duced to metallic antimony,	Yields 5 per cent. of water.	Stibico
	ing.	Yields 15 per cent. of water.	Volgeri
atom of		B. B. becomes magnetic, with borax gives the nickel reaction.	ZARA (Eme nick
lenable r		B. B. does not become magnetic. In O. F. colors the bead intensely violet (manganese).	DIALL (Rhot site)
nd a consta	Soluble in heated hydrochloric acid with effervescence (car-	phosphate of soda.	
- Poor	bonic acid).	B. B. becomes magnetic; with borax gives only the reactions for iron.	Siderit
niming of		Like Siderite, but after separation of the iron by ammonia gives a heavy precipitate with oxalate of ammonia.	ANKER
ut celati		Yields much water in matrass; does not become magnetic. Effervesces at first and then dissolves quietly.	Hydrot
Division 4. Division 4. Rearis or perfective soluble in Audrochloric or cuts and and additional or learning or consident of addres		Slowly soluble in HCl; the not too acid solu- tion gives with oxalic acid a white preci- pitate which on ignition becomes brick-red (oxide of cerium).	Parisite
1		Compare Smithsonite, Div. 2, p. 91. Streak yellow, usually crystalline. Water =10 per cent.	G öthit:
loric	Heated in R. F. become black and magnetic; quietly but	Streak yellow, not crystalline. Water = 144 per cent.	Limoni
sideroci,	difficultly soluble in HČl.	Streak red. Water = 5.3 per cent.	Turgite dro-her
۲ (Compare Hematite, Div. 2, page 70.	
huble	In the open tube give sulphur- ous acid. B. B. with soda	B. B. with soda deposits the brownish-red coating of oxide of cadmium.	Greeno
- All	give a sulphur reaction.	B. B. with soda on charcoal gives a coating of oxide of zinc.	Sphale: (Bler
Derfe	With borax give an amethys-	On charcoal with soda gives a coating of oxide of zinc.	Zincite
riv or	tine bead (manganese).	Gives much water in the closed tube.	WAD (I mang
	With borax give a deep blue bead (cobalt).	With soda on platinum wire gives a man- ganese reaction.	bolite
	With salt of phosphorus in O.F. give a yellow bead, which in	with nitrate of baryta.	Zippeit
	R.F. becomes deep green (uranium).	Gives no precipitate with nitrate of baryta.	URAND (Pitch l
	Colors the flame green, and when moistened with hydro- chloric acid colors the flame blue (copper).		

METALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture.	Lustre.	Hard- ness.	Sp. Gr.	Crystalliza tion.
u,Ćo,Ĺi², Ba, 湛l, 1,Ĥ².	Bluish-black.		Dull.	3.	8.2-8.6	
$=8b^{2}O^{3}+8b^{2}O^{5}$.	Yellowish.		Pearly.	4.5	4.08	IV.
+H20.	Yellow-red to white.		Pearly.	45.5	5.28	Mass.
⁵ +5aq.	White.	Pulverulent	Dull.	1	6.6	Mass.
0 ⁵ +6aq.	Emerald-green,		Vitreous.	33.25	2.6	
O ³ .	Rose-red, gray, brown.	Rhombohe- dral.	Vitreous. pearly.	8.5-4.5	3.5	ш.
'eC ⁵ O ⁹ .	Yellowish-white,gray, brown.	Rhombohe- dral.	Vitreous, pearly.	44.5	8.35	ш
) ³ . (Mn,Ca,Mg).	Ash-gray to brown- red.	Rhombohe- dral.	Vitreous.	3.5-4.5	3.7-3.9	III.
₹e,Mg)CO ³ .	White, gray, red.	Rhombohe- dral.	Vitreous,	3.5-4.	2.95-3.1	ш.
g,Ĥ²,Ċ.	White.	Basal.	Pearly.	2.	2.04	ш.
$\begin{array}{l} \text{La,Di}\text{CO}^{s} + \text{(Ca,}\\ \text{)}\text{F}^{2}\text{.} \end{array}$	Brown-yellow.	Basal.	Resinous.	4.5	4.35	ш,
04.	Dark-red, brown, black.	Prismatic.	Sub-metallic.	5.—5.5	4.3	IV.
² O ⁹ .	Brown-yellow, black.		Dull to sub- metallic.	5.	3.6-4.	
² O ¹ ,	Brown-black.		Dull to sub- metallic.	5.5	4.1-4.6	1
	Orange to honey-yel- low.	Prismatic.	Adamantine.	33.5	4.9	ш.
re)S.	White, yellow, green, brown, black,	Dodecahe- dral.	Resinous.	3.5-4.	3.9-4.2	L
(with MnO).	Orange-yellow to deep- red.		Adamantine.	44.5	5,68	ш.
1ºO5,	Gray, dull-black.		Dull.		1	
o,Ĉu,H².	Black.		Dull.	22.5	3.1-3.3	
)15+12aq.	Yellow.		Silky.	3.		
	Gray, brown, black.		Resinous.	5,5	6.4-8.	I,
O ¹¹ +5aq, contain- Ču.	Sky-blue to green.		Dull.	6.	2.6-2.8	

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MINERALS WITHOUT METALLIC LUSTRE

C. Infusible or fusible above 5.

DIVISION 4 (concluded).

Devinion 5 (in part).

II. MINERALS WIT

thout	General Characters.	Specific Characters,	Specie
uterto acid volthond Nica.	Moistened with sulphuric acid color the flame pale-green.		
(Continued.) hydrochloric acid or nitrio ieaving a residue of silica.		Fused with soda, the mass treated with water, and filtered, the residue dissolved in little HCl, the solution gives with oxalic acid a precipitate which ignited becomes	Monazite.
Jontin Judroc	ammonia (phosphoric acid).	After fusion becomes magnetic. Difficultly soluble in HCl.	Childreni
DIVERON 4.—(Continued.) or perfectly soluble is hydrochloric pelatintsing or leaving a re-	Fused with bisulphate of pot- assa, the mass dissolved in dilute hydrochloric acid and boiled with tin, gives a deep blue solution.	The dilute acid solution colors turmeric	Polycrase lymigni
D	With bisulphate of potassa, or	Gives reaction for the oxide of cerium. (See Monazite, above.)	Fluocerit
IN OF]	strong sulphuric acid, give the reaction for hydrofluorid	Evolves carbonic acid when treated with	Bastnäsit (Hamart
Nearly	acid.	Like fluocerite; but has an imperfect cleav- age in two directions.	Yttroceri
		With hydrochloric acid forms a perfect jelly. (Water = 11 per cent.)	
Noa.	of copper. Color yellow; after separation of the silica the solution	Decomposed without gelatinization. (Water = 20 per cent.)	Chrysoc
28		As above. (Water = 16 per cent.)	Cyanocha
DIVISION 5. Selatrites told hydrochtorto acid or are decomposed told the separation of silica. of B B (a the cheese inter weiver		Water = $12\frac{1}{2}$ per cent. In acicular crystals.	-
	Color white; massive; very hard.	After separation of the silica ammonia gives no precipitate, but oxalate of ammonia throws down oxalate of lime.	
	Gelatinize with hydrochloric acid.	The not too acid solution gives a precipitate with oxalic acid which becomes brick-red on ignition.	
DIVISION 5.		Does not gelatinize after ignition.	Thorite.
DIVISION			Wolchons ite.
loric ac			Genthite.
th hydroc	After long heating in R.F. be-	In the solution, after precipitation of the oxide of iron by ammonia, phosphate of soda gives a precipitate (magnesia).	
ize tolt		Gives only reactions for iron.	Chloropal
ann		Tompare Gillingite, Div. 5, p. 78.	
Ge	Moistened with cobalt solution	Gelatinizes with hydrochloric acid. Very light; absorbs water. B. B. shrivels up.	SEPIOLIT (Meerscha
	B.B. become pink.	Greasy feel ; does not adhere to the tongue.	CEROLITH

METALLIC LUSTRE.

Composition.	Color.	Cleavage or Fracture.	Lustre.	Hard- ness.	8p. Gr.	Oryscalli- sation.
² O ⁸ +Ca(Cl,F) ³ .	Colorless, white, blue, yellow, green.		Vitreous.	5.	8.2	III.
La,Di) ³ P ² O ⁸ +Th ²) ⁹ .	Yellow, clove-red, brown.	Basal.	Resinous.	5.—5.5	5.2	▼.
[n) ⁸ Al ² P ⁶ O ²⁹ +15ac	Yellow-brown to brownish-black.		Vitreous.	4.5-5.	8.18	IV.
ŕe,Ťi,Ĉb⁰, Ĥ⁰.	Black.		Sub-metallic.	5.5—6.5	4.8-5.1	IV.
·	Yellow, tile-red.		Weak.	45.	4.7 [.]	111.
'+Ce ² O ³ +4aq.	Wax-yellow.	Distinct.	Greasy.	4.	4.93	IV. ?
e,Y)F ² .	White, gray, blue.	[Weak, vit- reous.	4.5	3.45	
.SiO ⁴ .	Emerald-green.	Rhombohe- dral.	Vitreous.	5.	3.3	III.
	Blue to green.		Vitreous.	24.	2.2	
²,Ši,Ĥ².	Azure-blue.		Dull.	4.5	2.79	
⁶ Si⁵O ³⁰ +15ag.	Lemon-yellow.		Vitreous.		8 96	ſV.
iO ³ +H ² O.	White-gray.		-		2.71	
	Cherry-red, clove- brown.		Resinous.	5.5	4.9	
$0^4 + H^2 0.$	Orange, brown-black.		Resinous.	4.5-5.	5.—5.4	L
l,Fe,Mg,Si,H4.	Blue, grass-green.		Dull.	2.—2.5.	2.5	Amorph.
,Mg)'Si'O''.	Apple to emerald- green.		Resinous.	3.—4.	2.4	Amorph.
g, Fe, Ši, Ĥª.	Wood-brown to green.	Asbestiform	Glimmering.		2.4	Fib.
∂ ⁹ +5aq.	Pistachio-green to yel- low.		Earthy.	2.54.5	2.	Mass.
³ O ⁹ +2aq.	White, yellow, red.		 Dull.	2.—2.5	1.5	Maas.
² Si ² O ¹ +H ² O.	Green, yellow, white.	Conchoidal.	Resinous.	2 2.5		Mass.

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MINERALS WITHOUT METALLIC LUSTRE.

O. Infusible or fusible above 5.

DIVISION 5 (concluded).

DEVISION # (in part).

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II. MINERALS WIT:

	B. in the closed tube give water.	General Characters.	Specific Characters.	Specie
Division 5.—(Continued.) Belatinuse with hydrochloric acid or are decomposed with the separation of situa.		Decomposed by hydrochloric acid without gelatinizing. Loses on ignition 12-13 per cent. water.	pact and apple-green; bastite is foliated	Serpentii
the se	ed.) B.		Micaceous, with flexible but not elastic lamina.	Penninite
mued.)	Continued.)	Decomposed like the preced-	Compare pro-chlorite, ripidolite and deles- site. p. 95.	
(Oond	a)-(06		Crystalline foliated.	Monradite (Pyroxe
5	a	ing, but give only a little water in closed tube.	Very soft, with a soapy-feel.	Neolite.
DIVISION 5.—(Continued.) I or are decomposed with		water in closed tube.	Pearly lustre; perfect cleavage in one direc- tion.	SEYBERT
DIV acid or	traces.	Decomposed by hydrochloric acid with the formation of a jelly.	B. B. swells up and often glows with a bright light; strongly heated becomes grayish- green.	Gadolinit
chlori	but		With salt of phosphorus gives the fluorine reaction.	Chondro
A Aydro	water or		Fusible in very thin splinters; does not swell.	Gehlenite
ator tota	give no v		Infusible.	Chrysoli (olivine
Gelatinė	closed tube gi		After precipitation of the iron by ammonia, gives a precipitate with oxalate of ammo- nia (lime). Fusibility=5.	Monticell
	alo	Decomposed by hydrochloric	IT Compare Roepperite. p. 78.	
	the			Forsterite
	B, B, in	tinous silica.	Compare Monradite, Neolite, and Sey- bertite above.	1
	6) I	Decomposed without forming a jelly.	Generally crystallizes in trapezohedrons.	LEUCITE.
ns.		Micaceous; foliæ elastic. Give little or no water in the closed tube. Soft. H=12.5.	Decomposed by strong sulphuric acid (optic axial angle not exceeding 5°).	property and
oisiaip			Not decomposed by strong sulphuric acid (optic axial angle 44°-78°).	Muscovi
on 6. foregoing divisions.	under 7.		120).	MARGAR
8			Decomposed by sulphuric acid (optic axial angle 3°-20°, rarely less than 5°).	Phlogopi
DIVISI 0 to the	a) Hardness		Decomposed by sulphuric acid (optic charac- ters like muscovite).	MARGA DITE
DIVE Not belonging to the	a)	<u>a. 1991 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 1997 - 19</u>	Like muscovite; when decomposed by soda the hydrochloric solution gives a precipi- tate with sulphuric acid (baryta).	Oellacher
Not		Gives little water in closed tube (not always foliated); has a greasy feel. Soft.	When foliated the foliæ are not elastic.	Talc.

C.--Infusible or fusible above 5

METALLIC LUSTRE.

Composition.	Color,	Cleavage or Fracture.	Lustre.	Hard- ness.	Sp. Gr.	Orystalliza- tion.
² O ⁷ +2aq, with a ill amount of Fe. frequently col- i green by iron, cel, or chromium.	shades of oil-green, apple-green, yellow, red and black	Tough.	Sub-resinous, greasy, pearly, res- inous, silky, and earthy.	2.5—5.5	2.5—2.65	Only found in pseudo- morphs.
1Si ³ O ¹⁴ +4aq.	Green, gray, red.	Foliated.	Pearly.	2.5	2.7	ш.
Fe)SiO ³ -+H ² O.	Yellow.		Vitreous.	6.	8.27	Granular
I,Ši,H².	Green.		Silky.	12.	2.77	Fibrous.
Ca ⁶ , A l ² , F e ²)SiO ⁸ .	Yellow, copper-red, reddish-brown.	Foliated.	Pearly, sub- metallic.	4.—5.	33.1	IV.
,Be,Fe) ^s SiO⁵.	Blackish-green to black.		Vitreous.	6.5—7.	4.—4.5	IV.
i ³ O ¹⁴ .	White, red, yellow, brown, green.		Vitreous res- inous.	6.5	3.2	¦ ₩.
l F e)Si ² O ¹⁰ .	Gray-white.		Resinous.	5. 5—6.	8.	п.
^r e) ² SiO ⁴ .	Olive-green.	Prismatic.	Vitreous.	7.	8.8	īv.
[g) ² SiO ⁴ .	White-gray.	Prismatic.	Vitreous.	5.—5.5	3.—3.2	τ ν .
iO4.	White-gray.	Prismatic.	Vitreous.	6.—7.	3.2-3.3	IV.
	White-gray.		Vitreous.	5.5	2.4	
,Mg) ¹ Al ¹ Si ¹ O ²³ .	Green-black.	Foliated.	Splendent.	2.5	8.	ш.
Si ² O ⁸ .	White, gray, brown, green, yellow, red.	Foliated.	Pearly.	2.5	8.	I V .
$Si^{2}O^{11} + H^{2}O.$	White, gray, yellow, pink.	Foliated.	Pearly.	8.5-4.5	2.99	IV.
⁶ AlSi ⁵ O ²⁰ .	Yellow, red, white.	Foliated.	Pearly.	2.5	2.8	I V .
5i O*(+aq).	White-gray.	Foliated.	Pearly.	2.5	2.8	IV.
,Mg). Al²Si ⁴O¹ ⁴.	White, gray.	Foliated.	Pearly.		2.9	1
. ² Śi4O ¹⁹ .	White, apple to dark- green.	Foliated, compact.	Pearly.	 1.	8.7	IV.

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MINERALS WITHOUT METALLIC LUSTRE.

C. Infusible or fusible above 5.

DEVISION 6 (continued).

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			General Characters,	Specific Characters.		
				B. B. whitens and fuses on edges to a gray- yellow enamel.	B	
				Becomes black and magnetic.	P	
	not elastic. Give much water		Ï			
			posed by sulphuric acid.	Like ripidolite.	Í	
			, ,	Like ripidolite. Often gives reactions for chromium.	F	
				Easily distinguished by its hardness.	0	
			an emerald-green bead.	chromite, Div. 3, p. 71.		
			With soda on coal reduced to metallic tin.	Its high specific gravity is very noticeable.	C	
				Moistened with sulphuric acid colors the flame green (boric acid).	v	
	÷.	2	or with bisulphate of potash,	Prismatic cleavage.	1	
3	livielon		and boiled with tin, the solu- tion becomes violet (titanic scid). T Compare Scheelite,	- Octahedral cleavage.		
inned	otng (dnew		No cleavage.		
Cont	oreg	Hat		Compare perofskite, Div. 3, p. 71.	ľ	
о н 6. —С	to the f	inved.)	Fused with carbonate of soda,	The residue from the solution in water dis- solved in HCl colors turmeric paper orange- vellow (zirconia) B. B. swells up	1	
IVIBIO	onging	-(Cont	acid, and boiled with tin, the		1	
н	ot bel	ę.	tanic acid).	Found in octahedrons,	I	
	N		With soda fuses with effer-	silica is precipitated by addition of suffi-	- 0	
			Moistened with sulphuric acid colors the flame light-green.	Difficultly soluble in phosphorus salt to a colorless glass.	2	
		With phosphorus salt gives in O. F. a colorless bead, which	Decomposed by nitric acid, leaving a yellow residue of tungstic acid, which is soluble in alkalies.	2		
		Soluble in alkalies; not affected by nitric acid; occurs in soft earthy masses.				
			Amorphous. Gives much water in closed tube.	Soluble in sulphuric acid.]	
				Compare Kaolinite. Div. 1, p. 89.	-	
				vage surfaces show pearly lustre.	1	
		Cleaváble.	Cleavable in two directions, 1241°; fainter lustre than enstatite			
				Much like enstatite; on chargoal yields a magnetic mass.		
	DIVISION 6. – (Conténuel.)	DIVIZION 6. – (Continued.) Not belonging to the foregoing divisions.	Division 6 (Continued.) Not belonging to the foregoing divisions. a) (Continued.) Hardness under 7.	 not elastic. Give much water in the closed tube. Decom- posed by sulphuric acid. With salt of phosphorus give an emerald-green bead. With soda on coal reduced to metallic tin. Fused with carbonate of soda or with bisulphate of potash, dissolved in hydrochloric acid and boiled with tin, the solu- tion becomes violet (titanic acid). (S Compare Scheelite, p. 87. Fused with carbonate of soda, dissolved in hydrochloric acid, and boiled with tin, the solution becomes violet (ti- tanic acid). Gives water in the matrass. With soda fuses with effer- vescence to a clear glass. Moistened with sulphuric acid colors the flame light-green. Compare Lazulite, p. 90. With phosphorus salt gives in 0. F. a colorless bead, which in R. F., or better with tin on charcoal, becomes blue (when cold). Amorphous. Gives much water in closed tube. Cleavable. 	 Micaceous, the folis usually not elastic. Give much water in the closed tube. Decomposed by hydrochlorio acid. Micaceous, the folis usually not elastic. Give much water in the closed tube. Decomposed by hydrochlorio acid. Mith salt of phosphorus give by an emerald-green bead. With soda on coal reduced to metallic tim. With soda on coal reduced to metallic tim. Fused with carbonate of soda or with bisulphate of potash. dissolved in hydrochloric acid and boiled with tin, the solation in hydrochloric acid and boiled with tim, the solation becomes violet (titanic acid). Fused with carbonate of soda. Solve in hydrochloric acid and boiled with tim, the solution becomes violet (titanic acid). Fused with carbonate of soda. Solve in hydrochloric acid and boiled with tim, the solution becomes violet (titanic acid). Fused with carbonate of soda. Solve in hydrochloric acid and boiled with tim, the solution becomes violet (titanic acid). Fused with carbonate of soda. Solve in hydrochloric acid and boiled with tim, the solution in becomes violet (titanic acid). Fused with carbonate of soda. Solve in hydrochloric acid and boiled with tim, the solution in becomes violet (titanic acid). Fused with sulphuric acid colors the film light-green (borid colors the film light-green (borid colors the film light-green (borid of ammonium. Mistened with sulphuric acid colors bead, which in R. F. or better with tim on charcoal, becomes blue (when cold). Greenwable. Cleavable. 	

ETALLIC LUSTRE.

Composition.	Color.	Cleavage or Fructure.	Lustre.	Hard- ness.	Sp. Gr.	Crystalli- tation.
1:1Si ³ O ¹⁴ +4aq.	Shades of green to red.	Foliated.	Splendent, pearly.	2.5	2.7	v .
?e,Mg) ¹⁰ Al ³ Si ⁶ O ⁴² .	Green-black.	Foliated.	Pearly.	1.—2.	2.7—2.9	III. •
e, Hl, Fe, Si, H.	Dark olive-green.			12.5	2.89	
	White.	Basal.	Pearly.	2.5	2.65	111.
$c1Si^{3}O^{14}+4aq.$	Green, gray, red.	Basal.	Pearly.	2.5	2.7	111.
$ \begin{array}{l} Mg)(\Lambda l, Fe)SiO^{6} + \\ O. \end{array} $	Gray, green, black.	Basal.	Vitreous.	56.	3.5-3.6	V. ?
	Brown-black.		Adamantine and dull.	6.—7.	6.4-7.1	11.
ΓiO ³ +Mg ⁴ B ⁶ O ¹³ .	Brown-black.	Prismatic.	Sub-metallic.	34.	3.4	V. ?
	Red, brown, yellow, black.	Prismatic.	Adamantine.	6.5	4.2	11.
	Blue, brown, red, black.	Octahedral.	Adamantine.	5.5	8.8-3.9	11.
	Yellow, red, brown, black.		Adamantine.	5.5—6.	3.9-4.2	V
.a.Di,Fe,Y) ³ Cb ² ('fi) ³ O ¹⁴ .	Black.		Resinous.	56.	5.1	IV.
e,U) ⁸ Ti ² Cb ² O ¹² H ² O	Brown-black.		Brilliant.	6.5	4.9	IV.
ìi, Ťh, Ôe, Ôa, Fe, ι², F.	Brown-red.		Vitreous.	5.5	4.8	
+aq.	Colorless, milk-white, yellow, brown, red.		Vitreous.	6.—6.5	2.—2.8	Amorph
e)³₽³ O ³.	Yellow, brown, red.	Prismatic.	Resinous.	4.—5.	4.5	п.
04.	White, brown, yellow, red.		Vitreous.	4.5-5.	6.	11.
	Yellow.		Dull.			
e)O ³ +2aq.	White, brown, red.		Dull.		2.55	
Ю ³ .	White, gray, green, brown.	Prismatic.	Metalloidal.	5.5	8.2	IV.
Fe)SiO ³ .	Brown, gray, green.	Prismatic.	Sil ky .	5.5	8.2	IV.
Fe,SıO'.	Brown, green, black.	93°.	Metalloidal.	56.	3.39	IV.

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MINERALS WITHOUT METALLIC LUSTRE

C. Infusible or fusible above 5.

DIVISION 6 (concluded).

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MINERAL COAL.

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			General Characters.	Specific Characters.	Spe
			Compare Cassiterite, Rutile,	and opal which are very near 7 in hardness.	
			Pulverized and fused with bo- rax, colors the bead emerald- green (Chromium).	B. B. becomes blackish green, but cools to original color.	Ouva (chro net).
6	ons.		B. B., infusible and unaltered. With soda fuses, with effer- vescence to a clear glass (when pure).		Quarts
ΣI	plai	14	H = 7. Do not fuse to a clear	Difficultly fusible. $F_{\cdot} = 5 - 5.5$.	IOLITE
	med.)	o that	f = 1. Do not fuse to a clear glass with soda.	Infusible.	Staurc
-Infusible or Fusible above 5.	DIVISION 6(Continued.) Not belonging to the foregoing divisions.	Hardness 7 or more than		B. B. becomes colorless. Fused with soda, and the fusion dissolved in hydrochloric acid, the dilute acid solution colors turme- ric paper orange-yellow (zirconia).	1
	$ \vec{B}, \vec{B}, becomes milk white. Hexagonal with basal cleavage. $		B. B. becomes milk white. Hexagonal prisms,	Beryl.	
	B. B. becomes milk white. prisms, with right-angled clea	B. B. becomes milk white. Monoclinic prisms, with right-angled cleavage.	Euclas		
6	No	1		B. B. unchanged. Hexagonal prisms and pyramids, no basal cleavage.	Phenao
			H = 8. Gives with salt of phosphorus in open tube the fluorine reaction.	B. B. the yellow varieties become rose-red, crystallizes in prisms with perfect basal cleavage.	TOPAZ
			H = 7.5 - 8. Occurs generally in octahedrons.	With soda and borax on charcoal gives a coating of oxide of zinc.	GAHNI spin
				Soluble when pulverized in a bead of salt of phosphorus.	Spinel
		1	$\mathbf{H} = 10.$	Characterized by its hardness.	Diamo

MINE

FFT The native hydrocarbons are, for the most part, mixtures more analogous to rocks than true mineral species, and no atta

General Characters.	Specific Characters.	Va
Does not take fire in a lamp flame.	In closed tube yields a little water, and very little tarry product. B. B. burns with a feeble flame without fusing, leaving little ash; boiled with potash solution gives to it no color.	Anthr
Take fire in a lamp flame, and burn with a deep yellow flame, giving an empyreumatic odor.		Bitum
B. B. in glass tube give drops of tar or oil. Air dried Brown coal (Lignite) contains fre- quently from 15 to 20 p. c., or	Imparts little color to potash solution. The powder boiled with ether imparts to it a wine or brown-red color; very fusi- ble; flows in the flame of a candle like wax.	
more of water, which it loses when dried at 110° C.	Imparts to potash solution a brown color.	Brown

WUT METALLIC LUSTRE.

_	Composition.	Color.	Oleavage or Fracture.	Lustre.	Hard- Dors,	8p. Gr.	Crystallis tion.
	 Ca ^s &rSi ^s O ¹⁹ .	Emerald Green.		Vitreous.	7.5	3.5	I .
-	SiO ² .	Colorless, white, smoky, yellow, red, and all colors.	Conchoidal.	Vitreous.	7.	2.6	ш.
	(Mg,Fe) ² Al ² Si ⁵ O ¹⁸ .	Blue.		Vitreous.	7.	2.6	IV.
_	H ² (Mg,Fe) ² Al ⁶ Si ⁶ O ²⁴ .	Brown, red, black.		Vitreous-re- sinous.	7.	3.6	IV.
	ZrSiO'.	Colorless, red-gray, brown.		Adamantine	2.5	4.4-4.6	п.
-	Be ^s AlSi ⁶ O ¹⁸ .	Colorless, pink, blue- yellow and green.		Vitreous.	7.58.	2.6-2.7	III.
	H ² Be ² AlSi ² O ¹⁰ .	Mountain-green-blue, white.	Prismatic.	Vitreous.	7.5	8.1	. v.
		Colorless, yellow-red.	Conchoidal.	Vitreous.	7.5—8.	8.	III.
	A lSi(O, F²)⁵.	Colorless, white, yel- low, blue, pink.	Basal.	Vitreous.	8.	8.5	IV.
10	(Zn,Mg)(Al,Fe)O4.	Green, black.	Conchoidal.	Vitreous.	7.5.—8.	4.4-4.9	I.
	(Mg,Fe)(Al,Fe)O⁴.	Red, blue, green, yel- low, brown and black.	Conchoidal.	Vitreous.	8.	3.5-4.1	L.
-	С.	Colorless to black.	Octahedral.	Adamantine	10.	3.5-3.6	L

L COAL.

s made to classify them here, other than to state a few facts in regard to some varieties of mineral-coal, as given by Von Kobell.

	Composition.	Color.	Streak.	Lustre.	Hard- ness.	Sp. Gr.	
	C = 80—94 p. c.	Black.	Black.	Brilliant.	2.—2.5	1.8—1.7	
	50-85 p. c. residue on ignition.	Black.	Black.	Resinous.		1.2—1.8	
	С, Н, О.	Brown-black.	Brown.	Resinous.		1.—1.8	
•	Very variable.	Brown-black.	Brown.	Dull - resin- ous.			

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